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(54) Title: A NOVEL PLASTID-TARGETING NUCLEIC ACID SEQUENCE, A NOVEL β -AMYLASE SEQUENCE, A STIMULUS-RESPONSIVE PROMOTER AND USES THEREOF		
(57) Abstract		
The invention provides a novel chloroplast targeted novel β -amylase sequence (ct β -amylase), a novel chloroplast targeting nucleic acid sequence and a novel β -amylase sequence. There is also disclosed an inducible promoter which is independently stimulated by light or sugar stimulus. Methods of transforming plants using these sequences are described, as well as transformed plant cells, transformed plants and seed thereof, as well as chimaeric genes containing the sequences. Modification of starch levels in plants can be achieved, as well as the targeting of genes from the starch biosynthetic or degradative pathways, disease or pest resistance or variation of gene expression due to stimulus are described.		<p>The diagram illustrates the BNP10432 plasmid, which is 15001 bps long. Key features include:</p> <ul style="list-style-type: none"> 35Spro: Promoter region. ct bamY: Chloroplast targeting sequence for β-amylase. 35St: Promoter region. kan: Kanamycin resistance gene. npt: Neomycin phosphotransferase gene. colElori: Coliphage lambda endonuclease gene. RKori: Recombinant KpnI site. SacII,2902 / NotI,2904: SacII and NotI restriction sites. PstI,5312 / PvuII,5415: PstI and PvuII restriction sites. EcoRI,2 / 14997,SacI / 14989,NotI: EcoRI and SacII restriction sites. BamHI,14773 / EcoRI,14761 / SmaI,14756 / SalI,14751 / EcoRI,14684: BamHI, EcoRI, SmaI, SalI, and EcoRI restriction sites. PvuII,106 / PvuI,136 / BgIII,465: PvuII, PvuI, and BgIII restriction sites. 13578,EcoRI / 13545,PvuII: EcoRI and PvuII restriction sites. 12787,BamHI / 12781,PstI: BamHI and PstI restriction sites. 12347,HindIII / 12153,PvuII: HindIII and PvuII restriction sites. 11405,PvuI / 11283,ApaI: PvuI and ApaI restriction sites. 10454,PvuII / 10399,PstI / 10099,NheI / 10014,SacII / 9633,PvuII / 9014,BgIII: PvuII, PstI, NheI, SacII, PvuII, and BgIII restriction sites.

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A novel plastid-targeting nucleic acid sequence,
a novel β-amylase sequence, a stimulus-responsive promoter and
uses thereof

The precise mechanisms by which starch is synthesised and degraded in plants are unknown, despite the isolation and characterisation of a number of enzymes that are presumed to be involved in the process.

Starch is accumulated in the chloroplasts of leaves during the day and is used to supply the needs of the plant for energy and biosynthesis during the night. The mode by which this so-called transient starch is mobilised is not fully understood, but must involve the co-ordinated regulation of synthetic and degradative enzyme activities. In leaf tissues the main degradation pathway is thought to involve phosphorolytic and hydrolytic activities, especially α-glucosidase (E.C. 3.2.1.3) (Nielson and Stitt, 1997).

Starch is also accumulated in the amyloplasts in storage organs such as seeds, fruit and tubers. In this case starch is stored over longer periods of time and mobilisation of the starch is accompanied by degeneration of the storage organ tissues and increases in amylolytic and phosphorolytic activities. However, there is evidence to suggest that turnover of starch is also occurring in the amyloplasts of the storage organ (Sweetlove *et al*, 1996). This again requires the co-

ordinated regulation of the synthetic and degradative enzyme activities.

Chloroplasts and amyloplasts are both derived from proplastids and therefore have many characteristics in common besides being the site of starch synthesis in leaves and storage organs respectively; chloroplasts can be converted to amyloplasts and other types of plastid (Thomson and Whatley, 1980).

Starch is a mixture of two polysaccharides: amylose which is a linear chain of glucosyl units linked by α -1,4-glycosidic bonds; and amylopectin which is made up of many linear chains of α -1,4-polyglucans which are joined together by α -1,6 glycosidic bonds.

Enzymes involved in the synthesis of starch are ADPG pyrophosphorylase (E.C. 2.7.7.21), starch synthase (E.C. 2.4.1.21) and branching enzyme (E.C. 2.4.1.18). ADPG pyrophosphorylase is responsible for supplying the substrate ADPG, this molecule serving as the donor of glucose monomers which are linked together by the concerted action of starch synthases (α -1,4 bonds) and branching enzymes (α -1,6 bonds).

It is thought that the insoluble, crystalline structure of starch grains is formed by the close packing of the extended helical, branched amylopectin molecules, with the linear amylose molecules filling any spaces.

A range of starch-degrading enzyme activities has been

reported including α -amylase (E.C. 3.2.1.1), isoamylase (E.C. 3.2.1.68), β -amylase (E.C. 3.2.1.2), α -glucosidase (E.C. 3.2.1.3), starch phosphorylase (E.C. 2.4.1.1) and disproportionating enzyme (E.C. 2.4.1.25). Many of these enzyme activities exist in multiple forms in plants and some are thought to be involved in the synthesis of starch. All probably take part, to some extent, in the starch mobilisation process, however their exact roles and possible interactions are yet to be determined. The difficulties in attributing roles for the different enzymes is best exemplified by reference to two of the enzyme activities which are thought to be the major contributors to starch breakdown in plants: starch phosphorylase and amylase.

Starch phosphorylase catalyses the reversible release of glucose-1-phosphate from α -1,4-glucans. Two forms of starch phosphorylase are found in plant tissues: Pho1, or the L-type, is located inside plastids and has a high affinity towards maltodextrins; Pho2, or the H-type, is cytosolic and has high affinity to large, highly branched polyglucans such as glycogen. Although the plastidic Pho1 enzyme would be a likely candidate to be involved in the mobilisation of starch, antisense inhibition of the leaf enzyme activity had no effect on the starch accumulation in leaves of transgenic potato plants (Sonnewald et al., 1995). In another study, antisense inhibition of the cytoplasmic Pho2 had an influence on the

sprouting behaviour of transgenic potato tubers, but had no effect on the starch accumulation and degradation (Duwenig et al., 1997).

There are two major groups of amylase both of which hydrolyse α -1,4-glucosidic linkages in amylose and amylopectin: α -amylase acts randomly on non-terminal linkages, whereas β -amylase acts to release maltose units starting from the non-reducing end of the polyglucan chain. The subcellular location of α -amylase in the apoplastic space of plant cells is thought to reflect the fact that the enzyme is normally secreted. However, in a number of plants such as rice (Chen, et al., 1994) and sugar beet (Li, et al., 1992) the enzyme is also located inside chloroplasts and amyloplasts, despite the finding that the signal sequences at the amino-terminus of a number of α -amylase proteins are characteristic for translocation of protein across the ER membrane rather than the plastid membrane (Chen et al, 1994). In a study where the promoter and signal sequence of a rice α -amylase gene was fused to the bacterial GUS gene and introduced into rice, tobacco and potato using *Agrobacterium*-mediated transformation (Chan et al., 1994), it was demonstrated that the expressed GUS fusion protein was first transported to the endoplasmic reticulum and then exported into the culture medium of suspension cultures made from transgenic cells. It has been shown in a number of

studies that α -amylase will degrade native starch molecules.

In contrast, *in vitro* studies have shown that β -amylase will not degrade native starch granules without prior digestion of the granule with other enzymes. Mutants of rye (Daussant et al., 1981) and soybean (Hildebrand and Hymowitz, 1981) that lack active β -amylase or contain only traces of activity, respectively, apparently show normal growth and development. In addition, transgenic *Arabidopsis* plants in which the levels of β -amylase have been greatly reduced, do not show severe growth defects (Mita et al., 1997). Attempts to define the precise physiological role of β -amylases in plants have been hampered by inconclusive data concerning subcellular location. Although one study (Kakefuda et al., 1986) reported the presence of two β -amylases in pea chloroplasts, most studies involving species such as *Vicia faba*, barley, wheat, soybean, sweet potato and pea have concluded that most, if not all, β -amylase activity is extrachloroplastic (Nakamura et al., 1991). This view is supported by the fact that all β -amylase genes cloned to date encode proteins that lack amino-terminal chloroplast transit peptide sequences.

In cereals, three types of β -amylase have been described: an endosperm-specific form that accumulates during caryopsis maturation; a form that is synthesised *de novo* in aleurone cells of rice and maize during germination (Wang et al., 1996;

1997); and a β -amylase which is ubiquitous in vegetative organs. In *Arabidopsis*, the ubiquitous form accounts for approximately 80% of the total starch-degrading activity of rosette leaves. In common with all other β -amylase genes cloned to date, the gene for the ubiquitous *Arabidopsis* β -amylase does not encode a protein with a subcellular targeting signal, thus the enzyme is likely to be located in the cytosol.

The findings from a number of studies that the degradative activities can be removed without an adverse effect on the viability of the plant, plus the subcellular location of starch degrading enzymes outside the plastid, is surprising. The apparent absence of a plastid-localised β -amylase activity is especially surprising in light of the fact that the expected major end-product of β -amylase activity, namely maltose, has been identified as a product of starch degradation in isolated chloroplasts (Peavey et al., 1977). More recently, it has been shown that both glucose and maltose are exported from isolated cauliflower bud amyloplasts during the process of starch mobilisation (Neuhaus et al., 1995).

The ability to manipulate the amount of starch in the plastids of leaves or storage organs would be of high benefit to various industrial processes which utilise plant starches. For example, in an attempt to increase the starch content of potato tubers, it has been shown previously that when *E. coli*

ADPG PPase *glgC16* is overexpressed in transgenic potato tubers, there is an increase in flux of carbon into starch but there is only a small increase in net accumulation of starch (Sweetlove et al., 1997). Analysis of enzyme activities in the overexpressing lines showed that, apart from the alteration in ADPG PPase, the activity of amylase, specifically β -amylase was also altered. This data suggests that the accumulation of starch in tubers overexpressing *glgC16* protein is prevented by the breakdown of the newly synthesised starch, i.e. the starch is being turned over.

In another example, the availability of starch during the malting process is closely correlated with the types and amounts of degradative enzyme activities in the plant, specifically the storage organs. An increase in the degradative capacity of the crop would make the malting of cereal grain or the conversion of starch from tubers, or other storage organs, to alcohol more efficient and productive.

The type of starch present in the storage organ depends on the forms and activities of the ADPG pyrophosphorylase, starch synthase, branching enzyme and the degradative enzymes present. The interactions between the various enzymes will also be important.

There is considerable interest in creating novel starches *in planta* as this will reduce the costs of processing and modification of the starch before use in a variety of

industries such as food, paper, pharmaceuticals, glue, oil and textiles. The following examples show how starch hydrolytic activity can be important in altering the structure of starch *in vivo*.

It has been shown that, in maize kernels, the *sugary1* mutation causes the absence of a debranching enzyme which hydrolyses α -1,6-glycosyl linkages of starch (James et al., 1995). The mutation results in the decreased concentration of amylopectin and accumulation of the highly branched glucopolysaccharide, phytoglycogen.

It has been shown that in pea, short oligosaccharide molecules, starting with maltose and adding successive glucose units up to maltoheptose, specifically stimulate the activity of granule bound starch synthase I (GBSSI) (Denyer et al., 1996) which is generally accepted to be the major enzyme responsible for the synthesis of amylose (e.g. van der Leij et al., 1991; Hylton et al., 1995; Ainsworth et al., 1993). The manipulation of GBSSI activity by controlling the supply of malto-oligosaccharides is the subject of a recent patent (WO 97/16554) and suggests that an increase in the concentration of malto-oligosaccharides, and thus an increase in the ratio of amylose to amylopectin in the starch, can be brought about by the introduction of degradative enzymes namely α -amylase, β -amylase, disproportionating enzyme, debranching enzyme and starch phosphorylase. Patent WO 97/16554 also states that genes

for plastidial isoforms of these enzymes have been cloned. However, as discussed above, no β -amylase genes isolated to date encode a β -amylase enzyme with a protein targeting sequence and, in addition, there is doubt that α -amylases are originally targeted to plastids (Chen et al., 1994; Chan et al., 1994). Later in WO 97/16554, reference is made to the engineering of a suitable β -amylase cDNA sequence to add a plastid targeting sequence.

In addition to the industrial uses for starch in the storage organs, the amount of starch in the leaf has significant importance for the agronomy of a crop. Starch is synthesised in the leaf during daylight from the carbon fixed during photosynthesis. The starch is stored in the chloroplast and is broken down at night to become a source of energy and intermediates for metabolism in the plant. By which mechanisms the source-sink relationship is controlled are unknown at present, however, it is clear that manipulation of the amount and availability of the starch in leaf plastids will have a profound influence on plant productivity (biomass and yield).

The amount of starch in the leaf will also be important for those crops where the leaf is the major plant commodity, for example tobacco. It is known that starch content has an influence on the eventual flavour of tobacco when smoked. Provision of a means to manipulate the level of starch in

tobacco leaves could be of interest to the tobacco industry.

We describe here, for the first time, the isolation of a cDNA encoding a novel β -amylase enzyme which is targeted to plastids (henceforth known as chloroplast targeted (ct) β -amylase), by a novel targeting sequence. The isolation of this entire coding sequence is surprising, as it has generally been thought that β -amylase would only take part in the hydrolysis of starch once smaller polyglucan fragments had been released, either by translocation or through breakdown of the membrane, from the plastid into the cytoplasm. Location of the enzyme in plastids opens up the unforeseen possibility that ct β -amylase is involved in the degradation of transient starch located in chloroplasts and storage starch located in amyloplasts.

The similarity of characteristics between chloroplasts and amyloplasts (Thomson and Whatley, 1980) is of relevance to the current invention, as it has been shown that the transit peptides from chloroplast-targeted polypeptides can import heterologous polypeptides into amyloplasts and vice versa. For example, the transit peptide from the maize granule bound starch synthase enzyme when fused to the *E. coli* β -glucuronidase (GUS) protein will import the GUS protein not only into amyloplasts but also into chloroplasts (Klosgen and Weil, 1991).

In addition, we show that expression of the *ct-Bmy* gene in

Arabidopsis and the expression of *ct-B_my* promoter:GUS fusions in transgenic tobacco can be regulated independently by both light and sucrose. This is surprising in view of the tightly coupled light and sugar induction responses of *ATB-Amy* of *Arabidopsis* (Mita et al, 1995).

The present invention provides a nucleic acid sequence known herein as SEQ. ID. No. 1 and being from 1-294 nucleotides and having therewithin a sequence capable of targeting a further coding sequence to a plant plastid, or sequences being at least 65% or more homologous with the disclosed sequence SEQ. ID. No. 1 and having the same targeting ability.

Preferably the nucleic acid sequence encodes about 94 and more preferably about 85 amino acid residues.

The present invention also provides a nucleic acid sequence known herein as SEQ. ID. No. 2 and being from 1-1642 nucleotides and having therewithin a sequence capable of encoding β -amylase, or sequences being at least 65% or more homologous with the disclosed sequence within SEQ. ID. No. 2 and having the same encoding ability.

The present invention also provides a nucleic acid sequence known herein as SEQ. ID. No. 3 and being from 1-1953 nucleotides and having therewithin a sequence capable of encoding chloroplast targeted β -amylase, or sequences being at least 65% or more homologous with the disclosed sequence within SEQ. ID. No. 3 and having the same encoding ability.

Homologous sequences also include those sequences which hybridise to SEQ. ID. No. 1, SEQ. ID. No. 2 or SEQ. ID. No. 3 under medium stringency conditions (washing at 2x SSC at 65°C).

Preferably the nucleic acid sequence is an mRNA or cDNA sequence, although it may be genomic DNA.

The present invention also provides a method of increasing or decreasing in a plant the activity of an enzyme in the pathway of starch biosynthesis or degradation, the method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence and a coding sequence for an enzyme in the starch biosynthetic or degradative pathway, and regenerating a plant having an altered genome.

The present invention also provides a method of targeting proteins or enzymes to a plant plastid, the method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence and a coding sequence for a protein or an enzyme, and regenerating a plant having an altered genome, the protein or enzyme being one or more in the pathway of the following group: lipid synthesis, photosynthesis, amino acid metabolism, nitrogen fixation, carbon fixation or synthesis of carbohydrate polymers; or being able to confer a characteristic to the plant, the characteristic being selected from one or more of the following group: herbicide resistance

and pest resistance, for example, including fungal, bacterial or viral resistance.

The present invention also provides plants having therein a chimaeric gene comprising a promoter, a nucleic acid coding sequence encoding the plastid targeting sequence, the sequence being capable of targeting a coding sequence of an enzyme in the starch biosynthetic or degradative pathway to a plant plastid, and a terminator.

The present invention further provides a nucleic acid sequence capable of directing expression of a product encoded by a coding sequence which is operably linked thereto, said nucleic acid sequence being known herein as SEQ.ID. No. 8, or being at least 65% homologous therewith and having substantially the same function thereas, and said nucleic acid sequence being responsive to stimulus, the level of expression of said product being variable in response to the stimulus applied to said nucleic acid sequence.

The present invention further provides a method of varying the level of expression of a product encoded by a coding sequence operably linked to a nucleic acid sequence capable of directing expression of said product in a plant, said method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence capable of directing expression of a product encoded by a coding sequence that is operably linked thereto, said

nucleic acid sequence having substantially the sequence of SEQ.ID. No. 8 or being at least 65% homologous therewith and having substantially the same function thereas, and being responsive to stimulus.

Preferably the stimulus is the presence or absence of light and/or varying levels of sugar. Alternatively the stimulus is a stimulus which is developmentally controlled.

Advantageously the sugar is one or more of sucrose or glucose.

Preferably the sugar is sucrose.

Advantageously the inducible promoter, or nucleic acid sequence capable of directing expression of said product in a plant, is operable under conditions when there is no light but sugar is present, or when there is no sugar but light is present. The tissue of a plant where no light but sugar is present may suitably be underground organs or sink organs. Underground organs may be, for example, tubers, rhizomes or roots, whereas other sink organs may be young leaves or seeds.

The tissue of a plant where no sugar but light is present may be older leaves (where no sugar is transported), flower parts or germinating seeds.

Constructs and chimaeric genes having the DNA structural features described above are also aspects of the invention.

Plant cells containing a chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence

hereinabove described and a nucleic acid coding sequence of an enzyme in the starch biosynthetic or degradative pathway, or a chimaeric gene comprising a nucleic acid sequence capable of directing expression of a further coding sequence, or a chimaeric gene comprising a nucleic acid sequence hereinabove described that is responsive to stimulus and a coding sequence, the level of expression of said coding sequence being variable in response to the stimulus applied to said nucleic acid sequence are also an aspect of this invention, as is the seed of the transformed plant containing one or more chimaeric genes according to the invention.

Advantageously the plastid targeting sequence is the sequence SEQ. ID. No. 1.

In a first aspect of the invention the above method may be used to alter the metabolism of a leaf such that starch is accumulated therein or mobilised therefrom, this process altering the source-sink relationships within the plant as a whole. Such may be achieved by providing the targeting sequence and a nucleic acid coding sequence of an enzyme in the starch biosynthesis or degradative pathway under the direction of a suitable promoter. Suitable promoter selection would result in plants with increased or decreased levels of starch in the leaves which might be useful, for example, in the tobacco industry; or alternatively would result in changes in yield of starch in various other plant tissues such as tubers,

fruit and roots following modification of the source-sink relationships of the plant.

In this embodiment of the invention a suitable promoter would direct expression of the plastid targeting sequence and the coding sequence of an enzyme in the starch biosynthetic or degradative pathway throughout the whole plant, so called constitutive expression, or specifically to the leaves. These changes will have a profound effect such that the starch content and/or the yield of the organs of the plant would be significantly altered.

A preferred promoter capable of directing expression throughout all plant tissues is the promoter taken from cauliflower mosaic virus 35S gene. For leaf expression, preferred promoters can be taken from the gene for the small subunit of ribulose bisphosphate carboxylase or the pea plastocyanin gene. One skilled in the art will recognise other suitable promoters both for constitutive expression and specific leaf expression such as the nopaline synthase promoter and the chlorophyll a/b binding protein promoter respectively.

The coding sequence, or parts thereof, for the enzyme in the starch biosynthetic or degradative pathway may be arranged in the normal reading frame direction, i.e. sense, or in the reverse reading frame direction, i.e. antisense. Up or down regulation of the activity of the enzyme in a plant using sense, antisense or cosuppression technology (the latter as

described by DNAP in their European Patents Nos. 0465572 and 0647715) may be used to achieve alteration in the starch of the plant.

In a second aspect of the invention the inventive method may also be used to alter the metabolism of starch in storage organs such that starch content is increased and/or the starch is provided in a suitable form as required for the purposes of particular industrial processes. Such processes including paper making; manufacture of pharmaceuticals, textiles, dyes and building products; provision of baking, dairy and snack food products; making canned, dried or instant foods; malting of grain and production of syrups and alcohol.

In the first or second aspect of the method the enzyme selected for use in the chimaeric gene of the methods may be one from the starch degradative pathway, i.e. a starch degrading enzyme. Advantageously, the chimaeric gene comprises a chloroplast targeted β -amylase (hereinafter known as ct β -amylase), and more preferably comprises ct β -amylase derived from *Arabidopsis thaliana*, (hereinafter known as At ct β -amylase), see SEQ. ID. No. 3. Sequences homologous to At ct β -amylase which may be derivable from other plant sources such as potato, tobacco, wheat, maize and barley may also be used. Standard methods of cloning by hybridisation or polymerase chain reaction (PCR) techniques may be used to isolate

sequences from such organisms: for example molecular cloning techniques such as those described by Sambrook *et al.* (1989) and the PCR techniques described by Innes *et al.* (1990). Other starch degrading enzymes, the coding sequence of one or more of which would be suitable for use with the plastid targeting sequence, include α -amylase, disproportionating enzyme, debranching enzyme, starch phosphorylase, α -glucosidase and non-plastidic β -amylase.

In the second aspect of the inventive method preferred promoters which would direct expression to the storage organs of plants could be selected, for example, from the genes from the following list: the gene for high molecular weight glutenin of wheat endosperm; the gene for α,β -gliadin of wheat endosperm; the hordein gene of barley endosperm; or the gene for patatin from potato tubers. Other suitable promoters are known to those skilled in the art.

In either aspect of the invention, the alteration of tissue metabolism or alteration of starch type or characteristics may be made stimulus responsive, i.e. inducible, by virtue of use of the inducible promoter described herein (SEQ. ID. No. 8). For example, the light inducibility aspect of the inducible promoter could be used to manipulate seed set by inducing a gene such as Barnase (as exemplified in Patent WO 98/10081) to affect pollen development, or to affect

non-light responsive genes in otherwise light-dependant processes such as fruit ripening or seed germination. The light inducible promoter could also be used to turn on genes which affect secondary metabolite production in leaves, for example alkaloid production. Light inducible promoters may also be used to manipulate starch biosynthetic enzyme genes in leaves or other photosynthetic tissue, or for example in turning on genes after removal of tubers, for example, from storage in darkness. The sugar-inducibility aspect of the inducible promoter could be used to regulate genes in, for example, developing tuber or other non-photosynthetic tissue such as genes for pest resistance and/or genes which might affect the quality of the post-harvest crop. For potatoes, resistance genes to blight, blackleg and dry rot would be particularly of benefit and could be most advantageously cloned into recombinant genes with sugar inducible promoters. Alternatively, the sugar inducibility aspect of the inducible promoter could be used to drive the expression of genes for selectable markers in the tissue culture process.

One skilled in the art can readily delineate the sugar inducible responsive element from SEQ. ID. No. 8 and/or the light inducible responsive element by using well known techniques, such as deletion studies. Pwee and Gray (1993) describe such a deletion study within the pea plastocyanin gene using a marker gene in order to determine operative regions

thereof.

Methods described herein or in, for example, laboratory manuals by Sambrook et al (1989) and Gelvin and Stanton (1995) for cloning gene sequences and inserting them into appropriate carriers (vectors or plasmids etc.) are techniques well known to the skilled man for putting such concepts into effect. The chimaeric gene or genes as described above may be introduced on their own, or be accompanied by one or more other chimaeric genes, such as one or more of the other genes described above. In the case of the above described embodiments utilising a first chimaeric gene encoding an enzyme of the starch degradative pathway, the second chimaeric gene may, for example, comprise a nucleic acid sequence encoding an enzyme from the starch biosynthetic pathway also under the direction of a suitable promoter and a suitable terminator. The promoter and/or terminator of the second chimaeric gene may be the same as or different from the promoter and/or terminator of the first chimaeric gene. Suitable sequences encoding enzymes from the starch biosynthetic pathway are the nucleic acid sequences for sucrose synthase, ADPG pyrophosphorylase, starch synthase, and may also include branching enzyme, α -amylase, isoamylase, non-plastidic β -amylase, α -glucosidase, starch phosphorylase and disproportionating enzyme.

Methods for the introduction of more than one chimaeric gene into a plant have been described and comprise the

construction of a binary vector with the chimaeric genes joined together in one nucleic acid molecule; cotransformation using two or more different *Agrobacterium* cells, for example, with different binary vectors containing different chimaeric genes therein; or the transformation of a plant which already has a chimaeric gene with a second, different chimaeric gene, i.e. retransformation. In the latter case, the method of selection of transgenic plants after the introduction of the second chimaeric gene must be different from the selection method used for the introduction of the first chimaeric gene. Suitable selectable markers would include those for hygromycin, kanamycin, sulphonamide and Basta resistance. Biological methods such as crossing two plants, each plant containing a single chimaeric gene can also be used.

Use of two chimaeric gene constructs could be made in order to alter the starch content of an already transformed plant which shows a significant increase in a first enzyme activity and a consequent change in the synthesis of starch.

Thus, the present invention further provides a method of altering in a transgenic plant, which plant already shows an increase or decrease in an enzyme activity as a result of genetic transformation, a further enzyme in order to up or down regulate said further enzyme and thereby increase or decrease the amount of starch produced by the retransformed plant.

Advantageously the first transformed plant is a plant

having an increased enzyme activity in the starch biosynthetic pathway. An example of an attempt to increase the starch content of a plant is a transgenic potato transformed with the gene for ADPG-PPase, for example *glgC16* (see for example, WO 91/19806). The amount of starch increase in such a plant has been relatively small. This first transformed plant is advantageously retransformed with a chimaeric gene for a starch degrading enzyme, suitably comprising, for example, At ct β -amylase. The *glgC16* protein is expressed in the first transformed tubers and results in increased ADPG-PPase activity and an increase in flux of carbon to starch. Advantageously, the expression of the chimaeric At ct β -amylase gene, or parts thereof, in the retransformed tubers results in down regulation of the ct β -amylase activity, i.e. cosuppression or antisense technology, thus providing for an increase in starch accumulation.

Preferably the expression of the second enzyme is directed to tubers. A suitable promoter to direct the expression of the At ct β -amylase chimaeric gene in tubers is the promoter from the gene for patatin.

The first transformed potato plant expressing *glgC16* is kanamycin resistant, therefore the binary vector construct for the At ct β -amylase chimaeric gene carries a different resistance gene, suitably a gene for sulphonamide resistance,

for example. Increased starch production in the potato tuber would be of benefit, for example, to the potato crisp manufacturer as a 1% increase in potato dry matter would result in a 4% increase in product.

Potato crisp manufacture also serves to illustrate another benefit of the invention. When potato tubers are stored at temperatures below 8°C, reducing sugars, glucose and fructose from the breakdown of starch accumulate. When the potatoes are fried for crisps the reducing sugars react with amino acid in the Maillard reaction to give rise to brown colouration and off-tastes in the product. Introduction into potato plants of a chimaeric gene which would stop the breakdown of starch and thus the accumulation of reducing sugars would be of benefit to the snack food industry. Preferably the chimaeric gene would comprise the coding sequence, or a part of the sequence, for ct β -amylase in a cosuppression or antisense construct, driven by a suitable promoter and terminator. A suitable promoter would be taken from the gene for patatin in potato tubers. Advantageously any of the other starch degrading enzymes mentioned above could also be used instead of the ct β -amylase.

The inducible promoter of SEQ. ID. No. 8 could also be used in the construct if co-ordinated expression in the developing leaf and in the developing tuber were required, as the patatin promoter is also sucrose inducible (Rocha-Sosa et al (1989). Similarly, the sequence for the chloroplast

targeting polypeptide of SEQ. ID. No. 1 could also be used with any other gene which lacked its own targeting sequence and which was required to be directed to plastids.

The above examples serve to illustrate the possible benefits of using the present invention. One skilled in the art will recognise that the combination of genes and the plants to which the invention could be applied is considerable.

Gene combinations preferably will include ct β -amylase with one or more of the genes for sucrose synthase, ADPG pyrophosphorylase, starch synthase, branching enzyme, α -amylase, isoamylase, non-plastidic β -amylase, α -glucosidase, starch phosphorylase and disproportionating enzyme, the sequences of which are known to the skilled man. Alternatively, the targeting sequence from ct β -amylase may be used with one or more of the above genes.

The list of plants which could be transformed preferably include potato, wheat, maize, barley, tomato, rice, pea, soybean, peanut, cassava, yam, banana and tobacco.

The invention will now be described, by way of example, with reference to an embodiment for isolation of the cDNA for ct β -amylase from *Arabidopsis thaliana* and for incorporating the cDNA into tobacco and potato plants. Examples are also given on the stimulus responsive promoter and its activity in transgenic plants.

In order that the invention may be readily carried into effect reference will now be made, by way of example, to the following diagrammatic drawings in which;

Figure 1 shows the results of radiolabelled *in vitro* import translation products sampled on SDS-PAGE gel followed by fluorography. Legend: Molecular weight markers (lane M); translation products (lane Tr); chloroplasts reisolated and thermolysin-treated after import incubation (lane C); stromal fraction (lane S); washed thylakoids (lane T); thermolysin-treated thylakoids (lane tT); inner envelope fraction (lane I); outer envelope fraction (lane O). Putative precursor (P), intermediate (I) and mature (M) forms of β -amylase respectively. KiloDaltons (K);

Figure 2 shows the effect of light and the effect of light and sugars on the expression of ct β -amylase transcript in *Arabidopsis thaliana* seedlings. Figure 2a shows Northern blot analysis of total RNAs of 5-week old *Arabidopsis* plants grown in soil and exposed to 2 days continuous light (L), 2 days continuous darkness (D), 2 followed by 3 days of continuous light (LL) or 2 days of darkness followed by 3 days of continuous light (DL). Figure 2b shows Northern blot analysis of total RNAs of 5-week old *Arabidopsis* plants, grown *in vitro*, which were transferred either into water and exposed to 3 days continuous light (WL); or into 5% sucrose and exposed to 3 days of darkness (SD) or 3 days of continuous light (SL);

or into 5% glucose and exposed to 3 days of darkness (GD) or 3 days of continuous light (GL). Northern blots were hybridised with a radiolabelled *ct-Bmy* cDNA insert and autoradiographed (upper panels). The corresponding ethidium bromide-stained formaldehyde-agarose gels are shown in the bottom panels;

Figure 3 shows the diagrammatic representation of the T-DNA of the chimaeric *ct* β-amylase promoter-GUS genes constructed in Example 3 below, in which NosP represents the nopaline synthase promoter; NosT represents the nopaline synthase terminator; BR is the right border inverted repeat and BL is the left border inverted repeat of the T-DNA of pBII101; NPTII represents the neomycin phosphotransferase II coding sequence; GUS represents the β-glucuronidase coding sequence. *ct* β-amylase promoter fragments are represented by hatched rectangles; the PCR amplified Xho I-Bam HI bridging fragments are represented by black rectangles;

Figure 4 shows the effect of light and sucrose on the GUS activity expressed from a *ct Bmy* promoter-GUS chimaeric gene in tobacco seedlings;

Figure 5 shows the plasmid map of donator vector pDV35S (SK)V;

Figure 6 shows the plasmid map of donator vector pDV02000;

Figure 7 shows the plasmid map of binary plasmid pBNP10431 where 35Sp represents the CaMV 35S promoter, *ct bam*y represents the full length *ct* β-amylase cDNA, 35St represents the CaMV 35S

terminator, RB represents the right border of the binary vector pBinPlus, colE1ori represents the colE1 origin bacterial replication, RKori represents the oriV origin of replication of the RK2 plasmid, nptIII represents the neomycin phosphotransferase gene for bacterial resistance to kanamycin, LB represents the left border sequence of the binary vector, and kan represents the plant neomycin phosphotransferase recombinant gene required for plant resistance to kanamycin;

Figure 8 shows the plasmid map of binary plasmid pBNP10432 where abbreviations are as for Figure 7;

Figure 9 shows the plasmid map of binary plasmid pBNP02431 where abbreviations are as for Figure 7 except that patp represents the patatin class I promoter from vector pDV02000 in Figure 6 and nost represents the nopaline synthase terminator; and

Figure 10 shows the plasmid map of binary plasmid pBNP02432 where abbreviations are as for Figure 9.

In the sequence listing:

SEQ. ID. No. 1 is the nucleic acid capable of targeting a coding sequence to a plant plastid, particularly a chloroplast;

SEQ. ID. No. 2 is the nucleic acid which encodes β -amylase;

SEQ. ID. No. 3 is the complete sequence of chloroplast targeted (ct) β -amylase;

SEQ. ID. Nos. 4 and 5 are primers used in the

amplification process of Example 3;

SEQ. ID. Nos. 6 and 7 are primers used in the amplification process of Example 4; and

SEQ.ID. NO. 8 is the nucleic acid which is stimulus responsive, particularly to light and/or sugar.

Example 1

Isolation and characterisation of *Arabidopsis thaliana* chloroplast targeted β -amylase

Sequencing of cDNA insert in pBmy81

A BLASTN database search of the nucleotide sequence of a 37kb *Arabidopsis* chromosome IV DNA fragment in cosmid G16599 (Bevan et al., 1998) revealed the presence of a gene sharing significant homology with the extrachloroplastic β -amylase of *Arabidopsis*, barley maize rice soybean and rice. The search also identified several 3' terminal EST sequences, one of which, EST 81E10T7 (Newman et al., 1995), hereafter referred to as pBmy81, was identical over approximately 300 nucleotides. Clone EST 81E10T7 was supplied by the *Arabidopsis* Biological Resource Center (ABRC) DNA Stock Center (Ohio University, USA). A nested set of *Bal31* deletion subclones, spanning the cDNA insert in pBmy81, were used as DNA templates in double stranded PCR cycle sequencing reactions using fluorescent dye-labelled universal primers. Sequencing reactions were analysed on an

Applied Biosystems Model 373A automated sequencer. The nucleotide sequence of the cDNA insert in pBmy81 is shown in SEQ. ID No. 3. The construct pBmy81 was deposited by Advanced Technologies (Cambridge) Limited of 210 Cambridge Science Park, Cambridge CB4 4WA under the Budapest Treaty on the International Recognition of the Deposit of Micro-Organisms for the purposes of Patent Procedure at the National Collection of Industrial and Marine Bacteria (NCIMB), 23 St. Machar Street, Aberdeen Scotland on 4 August 1998 under Accession No. NCIMB 40964.

Identification of a putative chloroplast targeting signal

The pBmy81 cDNA insert comprises 36 untranslated nucleotides at the 5' end, an open reading frame (ORF) that encodes a protein of 548 amino acids and a 3' untranslated region (UTR) of 232bp. The protein encoded by the pBmy81 cDNA insert has a predicted molecular weight of 61 kDa and shares high amino acid similarity with plant extrachloroplastic β -amylases from maize, rice, barley, soybean and sweet potato. However, the protein encoded by pBmy81 differs from all other β -amylases reported so far in that it contains a unique N-terminal extension possessing the characteristics of a chloroplast targeting signal i.e. a high content of serine (16%), threonine (10%) and positively charged amino acid

residues (15%) (Baier and Dietz, 1997). Three domains which are distinguishing features of chloroplast targeting signals (Schatz and Dobberstein, 1996) were identified in the signal sequence: an uncharged amino-terminal domain; a central domain rich in hydroxylated amino acids; and a carboxy-terminal domain with the potential to form an amphiphilic β -strand.

cDNA insert in pBmy81 encodes a chloroplast targeted β -amylase

Intact chloroplasts were isolated from 50-60 g of pea shoots (*Pisum sativum* L. var Feltham First) using Percoll step-gradients. Plant material was grown and chloroplasts isolated according to the method of Mould and Gray (1997a).

The pBmy81 plasmid was linearised by restriction digestion with *NotI* and was transcribed *in vitro* using T7 RNA polymerase. Radiolabelled precursor protein was synthesised in a wheat germ translation system, including ^{35}S -methionine and ^{35}S -cysteine from transcripts of the pBmy81 cDNA essentially as described by Mould and Gray (1997b).

Import of radiolabelled *in vitro* translation products was performed as described by Mould and Gray (1997b). After the import incubation, intact chloroplasts were treated with thermolysin (0.2 mg/ml final concentration in import buffer) for 30 min on ice and then the protease reaction was stopped by the addition of EDTA to 50mM in import buffer. Chloroplasts were re-isolated through a cushion of 40% Percoll in import

buffer and then washed in import buffer (Mould and Gray, 1997b). An aliquot (1/10) of the thermolysin-treated chloroplast sample was taken for analysis and the remainder was fractionated essentially as described by Schnell and Blobel (1993). Samples of thermolysin-treated chloroplasts, stromal fraction, thylakoids and thermolysin-treated thylakoids were quantified by SDS-PAGE followed by coomasie blue staining and scanning densitometry of stained protein bands (subunits of ribulose bisphosphate carboxylase and light harvesting complex proteins were used as standards). Equivalent amounts of these fractions (approximately equal to 2% of the chloroplasts recovered from the Percoll gradient), and 50% of the inner and outer envelope fractions recovered, were analysed by electrophoresis on a 10% polyacrylamide gel in the presence of SDS, followed by fluorography. Results (Figure 1) show that the major translation product (lane Tr) was approximately 58 kDa. When isolated, intact pea chloroplasts were incubated with the radiolabelled protein in the presence of ATP, polypeptides of approximately 50 kDa and 48 kDa were generated (lane C). The resistance of these polypeptides to degradation by exogenously added thermolysin, indicates that they are products of radiolabelled protein import. Fractionation of the intact thermolysin-treated chloroplasts into stroma, washed thylakoids, thermolysin-treated thylakoids, inner envelopes and outer envelopes, demonstrated that the two radiolabelled

polypeptides were located in the stromal fraction.

EXAMPLE 2

Sucrose and light induction of *Arabidopsis thaliana* ct β-
amylase gene.

To demonstrate the induction of ct b-amylase in light *Arabidopsis thaliana* ecotype Landsberg plants were grown in the green house under an 18 hour light, 6 hour dark regime at 18°C. After 5 weeks, two trays of seedlings were transferred to complete darkness and two trays of seedlings were grown in continuous light. After two days, one tray of dark-adapted seedlings and a tray of light grown seedlings were used for isolation of total RNAs, and the second tray of each were exposed to a further 3 days of continuous light.

For combined sucrose-light-dark treatments, seeds of Landsberg ecotype were surface sterilised, placed on MS agar medium containing 1% sucrose and grown in a culture room with an 18 hour light, 6 hour dark regime. Five week old seedlings were transferred onto sterilised distilled water or a 5% solution of sucrose or glucose in water. The seedlings were maintained either in continuous light or darkness for three days. Total RNAs were prepared from seedlings of each test and were analysed by northern blot analysis as described by Eggermont et al. (1996). Northerns were probed with the gel-purified cDNA insert in pBmy81 following random labelling with

³²P-dCTP as described by Feinberg and Vogelstein (1983).

The results shown in Figure 2A indicate that the ct β -amylase genetranscript is inducible due to light.

The results shown in Figure 2B indicate that the ct β -amylase transcript is induced in the dark with 5% sucrose and to a lesser extent with 5% glucose. This induction is enhanced further in the light in the presence of the sugars. These results show that the effect of light and sugars are independent of each other.

EXAMPLE 3

Construction of ct β -amylase promoter-GUS fusions

Promoter fragments were isolated from the ct β -amylase gene located in cosmid G16599 (Bevan et al., 1998) by restriction enzyme digestion. Convenient restriction sites in the promoter were *Hind* III at nucleotide position -1662 bp (starting at 19179bp on the minus strand of SEQ. ID. No. 8), *Sal* I at -1127 bp and *Pst* I at -371 bp and an *Xho* I site located at position +21 bp downstream of the ct β -amylase initiating methionine were used to isolate three different lengths of promoter plus transit peptide sequence (the A of the translation initiation methionine ATG is numbered +1).

A 294 bp (SEQ. ID. No. 1) fragment of the ct β -amylase gene located in cosmid G16599 (Bevan et al., 1998) was amplified using the oligonucleotide primers:

SEQ. ID. No. 4

P1: (5' - AAT TCC TCG AGT TCT CTT ATC - 3') and

SEQ. ID. No. 5

P2: (5' - cgg gAT CCC TGA CAT TGT TAC - 3').

In primer P1, the underlined bases refers to the *Xho* I site located at position +21 bp; in primer P2 the bases in lower case refer to the nucleotides added in order to create a *Bam* HI site.

Chimaeric ct β -amylase promoter-GUS genes were created by triple ligation of the promoter fragment; the PCR bridging fragment digested with *Xho* I and *Bam* HI; and the GUS vector pBI101 (Jefferson et al., 1987) digested with *Hind* III-*Bam* HI, *Sal* I-*Bam* HI or *Pst* I-*Bam* HI (Figure 3). Constructs were termed H β GUS, S β GUS and P β GUS respectively.

The chimaeric gene constructs were transferred to *Agrobacterium tumefaciens* LBA4404 by triparental mating (Bevan, 1984) and introduced into *Nicotiana tabacum* var Samsun by the leaf disk transformation method (Horsch et al., 1985).

EXAMPLE 3A

Sucrose and light induction of chimaeric *Arabidopsis thaliana*ct β -amylase promoter-GUS gene in tobacco seedlings.

Plants containing the H β GUS and P β GUS constructs expressed high levels of GUS activity and F1 seedling progeny of the lines were used to investigate light and sucrose inducible expression of the chimaeric genes. F1 tobacco seeds were surface sterilised, placed on MS agar medium containing 1% sucrose and grown in a culture room with an 18 hour light, 6 hour dark regime. Two to three week old seedlings were transferred onto a 5% sucrose solution or onto distilled water and maintained either in continuous light or darkness for three days. Total protein extracts from pools of 10 to 14 seedlings were analysed for GUS activity using the fluorogenic substrate 4-methylumbelliferyl-glucuronide (4-MUG) as described by Jefferson et al. (1987). With both constructs, the level of GUS activity in seedlings exposed to continuous light in the absence of sucrose was similar to the levels of GUS activity in seedlings exposed to sucrose in the absence of light (Figure 4). However, exposure of seedlings to both continuous light and sucrose increased levels of GUS activity by approximately two to three fold. These results are broadly in agreement with the results from the experiments with the ct b-amylase gene itself which showed that light inducibility and sucrose inducibility are independent processes.

Histochemical staining for GUS showed that activity was detected in the cotyledons of two week old seedlings and little or no activity in the first true leaves or in the stems and roots. In four week old seedlings, additional GUS activity was shown throughout the first true leaves and also in the stems. GUS staining was particularly associated with chloroplast-rich parenchyma (chlorenchyma) cells located between the xylem rays and between xylem and the bundles of phloem that constitute the internal phloem in stems.

EXAMPLE 4

Construction of ct β -amylase plasmids for use in transformation
of tobacco and potato leaves

Site-directed mutagenesis was used to convert the *Kpn* I site located at position 2302 bp of the pBmy81 plasmid to a *Bam* HI site. Oligonucleotide primers

SEQ. ID. No. 6

P3: (5'- GCT GGT ACG CCT GCA GGA TCC GGT CCG GAA TTC CC - 3')
and

SEQ. ID. No. 7

P4: (5'- GGG AAT TCC GGA CCG GAT CCT GCA GGC GTA CCA GC - 3')

were designed and used with the Quick Change site-directed

mutagenesis kit (Promega). Protocol was as outlined by the manufacturer.

The full length ct β -amylase coding sequence was excised from the mutated pBmy81 plasmid by cleavage with *Bam* HI and then purified with GeneClean (BIO 101). The *Bam* HI fragment was ligated into the *Bam* HI site of the donator vectors pDV35S(SK)V (see Figure 5) and pDV02000 (see Figure 6). pDV35S(SK)V consists of pBluescript (Stratagene) carrying a 35S CaMV promoter-35S terminator, similar constructs are known in the art (e.g. Odell et al., 1985). pDV02000 consists of pBluescript with a 1.4 kbp patatin promoter-nopaline synthase terminator. One skilled in the art could make similar constructs from known sequences (e.g. Liu et al., 1990). Plasmids with the coding sequence in both the sense and antisense orientation relative to the promoters were isolated, and the ct β -amylase chimaeric genes subcloned from the donator vectors into the binary vector pBinPlus (van Engelen et al., 1995). The plasmid maps are shown in Figures 7-10.

EXAMPLE 5

Transformation or Retransformation of plants

Potato plants were transformed using the method of leaf disk cocultivation as essentially described by Horsch (1985). The binary vectors as described above were transferred to

Agrobacterium tumefaciens LBA4404 using the method of electroporation, and cultures of said *Agrobacteria* used in transformation so that regenerated plants carry the chimaeric genes as described in Example 4.

The patatin promoter-ct β -amylase-nopaline synthase terminator chimaeric gene binary plasmid, can be used to transform a potato plant already carrying the chimaeric gene for *E. coli* ADPG-Ppase *glgC16* by the methods of leaf disk cocultivation.

EXAMPLE 6

Construction of plasmids with the targeting peptide of AT ct β -amylase

The plastid targeting sequence of AT ct β -amylase is contained within a 294 bp fragment equivalent to SEQ. ID. No. 1. PCR amplification or restriction enzyme digestion can be used to isolate fragments of DNA from the plasmids described in Example 3, i.e. fragments will consist of the 35S CaMV promoter plus plastid targeting sequence or the patatin promoter plus the plastid targeting sequence. Chimaeric genes can be constructed by ligating coding sequences for proteins or enzymes as translational fusions with the transit peptide sequence. Translated proteins would be transported to the plastids to provide novel activities or to affect metabolic pathways.

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INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

- A. The indications made below relate to the microorganism referred to in the description
on page 29, lines 9/10

B. IDENTIFICATION OF DEPOSIT

Further deposits are identified on an additional sheet

Name of depositary institution

The National Collections for Industrial and Marine Bacteria Limited (NCIMB)

Address of depositary institution (*including postal code and country*)

23 St. Machar Drive
Aberdeen AB2 1RY
Scotland, United Kingdom

Date of deposit	Accession Number
4 August 1998	NCIMB 40964

C. ADDITIONAL INDICATIONS (*leave blank if not applicable*)

This information is continued on additional sheet

Please find enclosed copies of the Receipt of Deposit and copies of the viability proofs from the Depositary Institution. Name and Address of Depositor: Advanced Technologies (Cambridge) Limited, 210 Cambridge Science Park, Cambridge CB4 0WA, United Kingdom.

- Characteristics: E. Coli XLI Blue MRF1
- Produced by: inserting 1953bp sequence for ct β-amylase from Arabidopsis thaliana ecotype Columbia inserted into a vector type puC type-pZL1 (Gibco BRL) which is in an E. coli strain XLI Blue MRF1
- Usefulness: targeting a particular β-amylase to a chloroplast

D. DESIGNATED STATES FOR WHICH INDICATIONS ARE MADE (*if the indications are not for all designated States*)E. SEPARATE FURNISHING OF INDICATIONS (*leave blank if not applicable*)

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FOR THE PURPOSES OF PATENT PROCEDURE**

<p>Advanced Technologies (Cambridge) Ltd., 210 Cambridge Science Park, Cambridge, CB4 4WA</p>	<p>INTERNATIONAL FORM RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT issued pursuant to Rule 7.1 by the INTERNATIONAL DEPOSITORY AUTHORITY identified at the bottom of this page</p>						
<p>NAME AND ADDRESS OF DEPOSITOR</p> <p>I. IDENTIFICATION OF THE MICROORGANISM</p> <table border="0" style="width: 100%; border-collapse: collapse;"> <tr> <td style="width: 50%; vertical-align: top; padding: 5px;">Identification reference given by the DEPOSITOR:</td> <td style="width: 50%; vertical-align: top; padding: 5px;">Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY:</td> </tr> <tr> <td style="padding: 5px;"><i>Escherichia coli</i> (XL1 Blue MRF pBmy81)</td> <td style="padding: 5px;">NCIMB 40964</td> </tr> </table> <p>II. SCIENTIFIC DESCRIPTION AND/OR PROPOSED TAXONOMIC DESIGNATION</p> <p>The microorganism identified under I above was accompanied by:</p> <p><input type="checkbox"/> a scientific description</p> <p><input checked="" type="checkbox"/> a proposed taxonomic designation</p> <p>(Mark with a cross where applicable)</p> <p>III. RECEIPT AND ACCEPTANCE</p> <p>This International Depository Authority accepts the microorganism identified under I above, which was received by it on 4 August 1998 (date of the original deposit)¹</p> <p>IV. RECEIPT OF REQUEST FOR CONVERSION</p> <p>The microorganism identified under I above was received by this International Depository Authority on (date of the original deposit) and a request to convert the original deposit to a deposit under the Budapest Treaty was received by it on (date of receipt of request for conversion)</p> <p>V. INTERNATIONAL DEPOSITORY AUTHORITY</p> <table border="0" style="width: 100%; border-collapse: collapse;"> <tr> <td style="width: 50%; vertical-align: top; padding: 5px;">Name: NCIMB Ltd., Address: 23 St Machar Drive, Aberdeen, AB24 3RY, Scotland.</td> <td style="width: 50%; vertical-align: top; padding: 5px;">Signature(s) of person(s) having the power to represent the International Depository Authority or of authorised official(s):  Date: 19 August 1998</td> </tr> </table>		Identification reference given by the DEPOSITOR:	Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY:	<i>Escherichia coli</i> (XL1 Blue MRF pBmy81)	NCIMB 40964	Name: NCIMB Ltd., Address: 23 St Machar Drive, Aberdeen, AB24 3RY, Scotland.	Signature(s) of person(s) having the power to represent the International Depository Authority or of authorised official(s):  Date: 19 August 1998
Identification reference given by the DEPOSITOR:	Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY:						
<i>Escherichia coli</i> (XL1 Blue MRF pBmy81)	NCIMB 40964						
Name: NCIMB Ltd., Address: 23 St Machar Drive, Aberdeen, AB24 3RY, Scotland.	Signature(s) of person(s) having the power to represent the International Depository Authority or of authorised official(s):  Date: 19 August 1998						

¹ Where Rule 6/4(d) applies, such date is the date on which the status of International Depository Authority was acquired.
Form BP/4 (sole page)

**BUDAPEST TREATY ON THE INTERNATIONAL
RECOGNITION OF THE DEPOSIT OF MICROORGANISMS
FOR THE PURPOSES OF PATENT PROCEDURE**

**Advanced Technologies (Cambridge) Ltd.,
210 Cambridge Science Park,
Cambridge.
CB4 4WA**

INTERNATIONAL FORM

VIABILITY STATEMENT

**Issued pursuant to Rule 10.2 by the
INTERNATIONAL DEPOSITORY AUTHORITY
Identified on the following page**

**NAME AND ADDRESS OF THE PARTY
TO WHOM THE VIABILITY STATEMENT
IS ISSUED**

I. DEPOSITOR		II. IDENTIFICATION OF THE MICROORGANISM
Name: AS ABOVE	Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY: NCIMB 40964	
Address:	Date of the deposit or of the transfer ¹ :	
	4 August 1998	
III. VIABILITY STATEMENT		
The viability of the microorganism identified under II above was tested on 8 August 1998		2. On that date, the said microorganism was:
3		
<input checked="" type="checkbox"/>	viable	
3		
<input type="checkbox"/>	no longer viable	

- 1 Indicate the date of the original deposit or, where a new deposit or a transfer has been made, the most recent relevant date (date of the new deposit or date of the transfer).
 - 2 In the cases referred to in Rule 10.2(a)(ii) and (iii), refer to the most recent viability test.
 - 3 Mark with a cross the applicable box.

IV. CONDITIONS UNDER WHICH THE VIABILITY TEST HAS BEEN PERFORMED⁴**V. INTERNATIONAL DEPOSITORY AUTHORITY**

Name: NCIMB Ltd.,

Address: 23 St Machar Drive,
Aberdeen,
A24 3RY,
Scotland.

Signature(s) of person(s) having the power
to represent the International Depository
Authority or of authorised official(s):



Date: 19 August 1998

⁴ Fill in if the information has been requested and if the results of the test were negative.

CLAIMS

1. A nucleic acid sequence known herein as SEQ. ID. No. 1 and being from 1-294 nucleotides and having therewithin a sequence capable of targeting a further coding sequence to a plant plastid, or sequences being at least 65% or more homologous with the disclosed sequence SEQ. ID. No. 1 and having the same targeting ability.
2. A nucleic acid sequence according to Claim 1, wherein said nucleic acid sequence encodes about 94 amino acid residues.
3. A nucleic acid sequence according to Claim 2, wherein said sequence encodes about 85 amino acid residues.
4. A nucleic acid sequence known herein as SEQ. ID. No. 2 and being from 1-1642 nucleotides and having therewithin a sequence capable of encoding β -amylase, or sequences being at least 65% or more homologous with the disclosed sequence within SEQ. ID. No. 2 and having the same encoding ability.
5. A nucleic acid sequence known herein as SEQ. ID. No. 3 and being from 1-1953 nucleotides and having therewithin a sequence capable of encoding chloroplast targeted β -amylase, or sequences being at least 65% or more homologous with the disclosed sequence within SEQ. ID. No. 3 and having the same encoding ability.

6. A nucleic acid sequence capable of directing expression of a product encoded by a coding sequence which is operably linked thereto, said nucleic acid sequence being known herein as SEQ. ID. No. 8, or being at least 65% homologous therewith and having substantially the same function thereas, and said nucleic acid sequence being responsive to stimulus, the level of expression of said product being variable in response to the stimulus applied to said nucleic acid sequence.
7. A nucleic acid sequence according to any one of the preceding claims, wherein said nucleic acid sequence is an mRNA, cDNA or a genomic DNA sequence.
8. A method of increasing or decreasing in a plant the activity of an enzyme in the pathway of starch biosynthesis or degradation, the method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence and a coding sequence for an enzyme in the starch biosynthetic or degradative pathway, and regenerating a plant having an altered genome.
9. A method according to Claim 8, wherein said nucleic acid sequence comprises the sequence of Claim 1.
10. A method according to Claim 8, wherein said enzyme in the starch degradative pathway is β -amylase.
11. A method according to Claim 8, 9 or 10, wherein said

coding sequence is the nucleic acid sequence of Claim 4, being a coding sequence for an enzyme in the starch degradation pathway.

12. A method according to Claim 8, wherein said nucleic acid sequence is the nucleic acid sequence of Claim 5.
13. A method according to any one of Claims 8-12, wherein said enzyme from the starch biosynthetic pathway is one of the group consisting of sucrose synthase, ADPG pyrophosphorylase, starch synthase, branching enzyme, α -amylase, isoamylase, non-plastidic β -amylase, α -glucosidase, starch phosphorylase and disproportionating enzyme.
14. A method of targeting proteins or enzymes to a plant plastid, the method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence and a coding sequence for a protein or an enzyme, and regenerating a plant having an altered genome, the protein or enzyme being one or more in the pathway of the following group: lipid synthesis, photosynthesis, amino acid metabolism, nitrogen fixation, carbon fixation or synthesis of carbohydrate polymers; or being able to confer a characteristic to the plant, the nucleic acid sequence being the nucleic acid sequence of Claim 1.

15. A method according to Claim 14, wherein said characteristics selected from one or more of the following group: herbicide resistance and pest resistance.
16. A method of varying the level of expression of a product encoded by a coding sequence operably linked to a nucleic acid sequence capable of directing expression of said product in a plant, said method comprising the steps of stably incorporating into a plant genome a chimaeric gene comprising a nucleic acid sequence capable of directing expression of a product encoded by a coding sequence that is operably linked thereto, said nucleic acid sequence having substantially the sequence of SEQ. ID. No. 8 or being at least 65% homologous therewith and having substantially the same function thereas, and being responsive to stimulus.
17. A method according to Claim 16, wherein said coding sequence encodes a gene causing sterility in said plant during flowering and which is caused to be expressed by said nucleic acid sequence upon exposure to light.
18. A method according to Claim 16, wherein said coding sequence encodes a gene which alters the level of starch biosynthetic enzyme genes in leaves or other photosynthetic tissue.
19. A method according to Claim 16, wherein said coding sequence encodes a gene whose expression level is varied

- upon the generation of sugar in a developing storage organ and the effect of sugar on said nucleic acid sequence.
20. A method according to Claim 16, wherein said nucleic acid sequence is operable when there is light but no sugar.
 21. A method according to Claim 16, wherein said nucleic acid sequence is operable when there is sugar but no light.
 22. A method of altering in a transgenic plant, which plant already shows an increase or decrease in an enzyme activity as a result of genetic transformation, a further enzyme in order to up or down regulate said further enzyme and thereby increase or decrease the amount of starch produced by the retransformed plant.
 23. A method according to Claim 22, wherein said first transformed plant is a plant having an increased or decreased enzyme activity in the starch biosynthetic pathway.
 24. A method according to Claim 23, wherein said first transformed plant is a transgenic potato transformed with the gene for ADPG-PPase.
 25. A method according to any one of Claims 8, 14 or 16, wherein said chimaeric gene further comprises a promoter selected from the group consisting of the cauliflower mosaic virus 35S promoter (full or truncated), the rubisco promoter, the pea plastocyanin promoter, the nopaline synthase promoter, the chlorophyll r/b binding promoter,

the high molecular weight glutenin promoter, the α , β -gliadin promoter, the hordein promoter and the patatin promoter.

26. A method according to any one of Claims 8, 14 or 16, wherein said coding sequence of said enzyme in said chimaeric gene provides up or down regulation of the activity of said enzyme.
27. A method according to Claim 16, wherein said stimulus is the presence or absence of light and/or varying levels of sugar.
28. A method according to Claim 16, wherein said stimulus is a stimulus which is developmentally controlled.
29. A method according to Claim 27, wherein the sugar is one or more of sucrose or glucose.
30. A method according to Claim 29, wherein the sugar is sucrose.
31. A chimaeric gene comprising a nucleic acid sequence according to Claim 1.
32. A chimaeric gene comprising a nucleic acid sequence encoding a plastid targeting sequence according to Claim 1 and a nucleic acid coding sequence of an enzyme in the starch, degradative pathway, the enzyme being encoded by the sequence according to Claim 4.
33. A chimaeric gene comprising a nucleic acid sequence capable of directing expression of a further coding

sequence, said coding sequence being the sequence according to Claim 4.

34. A chimaeric gene comprising a nucleic acid sequence according to Claim 6 that is responsive to stimulus and a coding sequence, the level of expression of said coding sequence being variable in response to the stimulus applied to said nucleic acid sequence.
35. Plant cells comprising the nucleic acid sequence of any one of Claims 1, 4, 5 or 6.
36. Seed of the transformed plant containing one or more of the nucleic acid sequences of Claims 1, 4, 5, or 6.
37. Plants transformed according to the methods of any one of Claims 8, 14 or 16, where in said plants are one or more of the group consisting of potato, wheat, maize, barley, tomato, rice, pea, soybean, peanut, cassava, yam, banana and tobacco.

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Fig.1.



Fig.2(a).

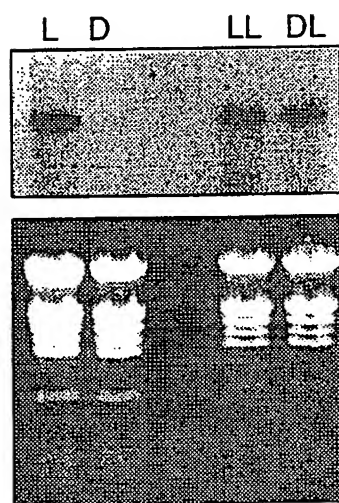


Fig.2(b).

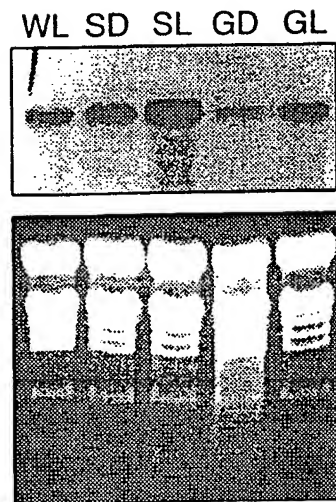
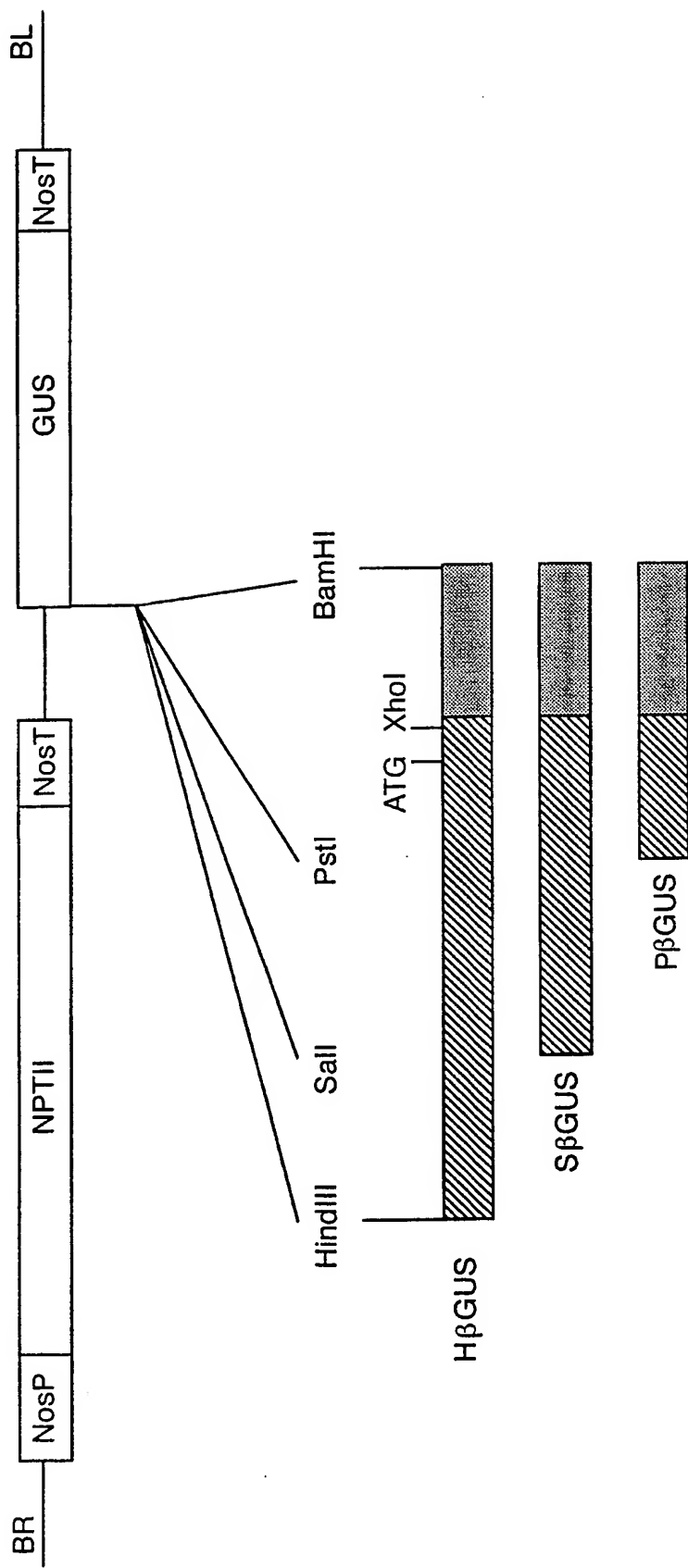


Fig.3.

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Fig.4.

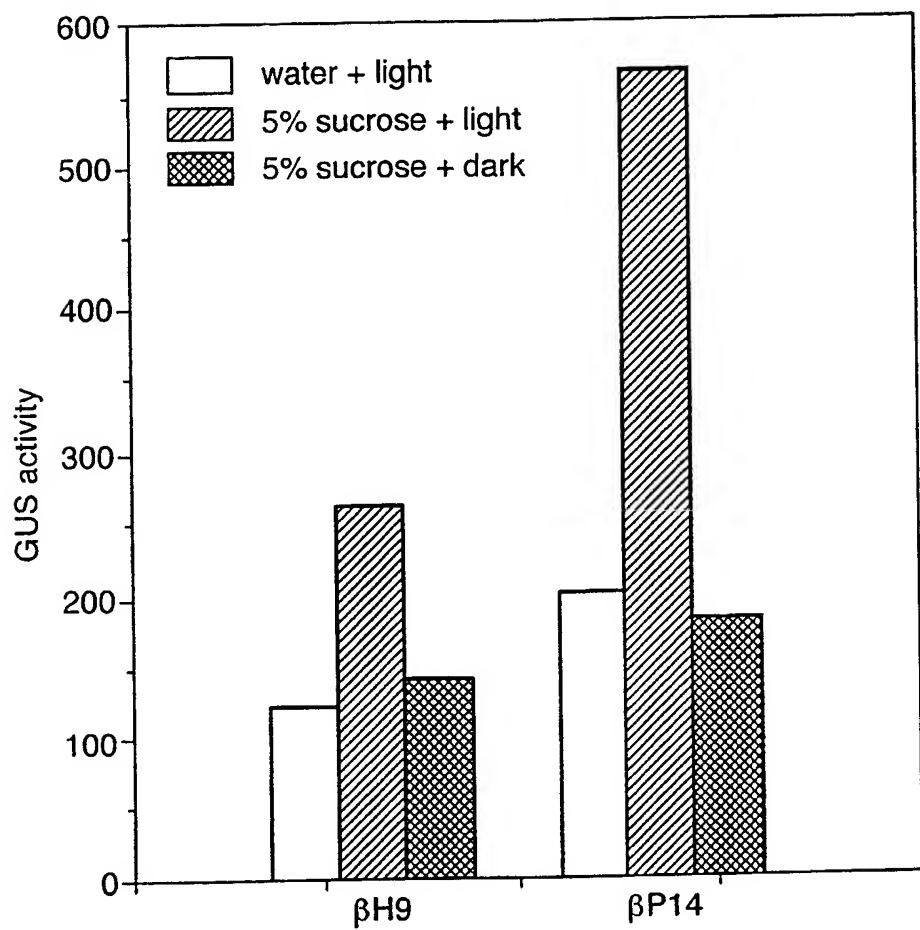


Fig.5.

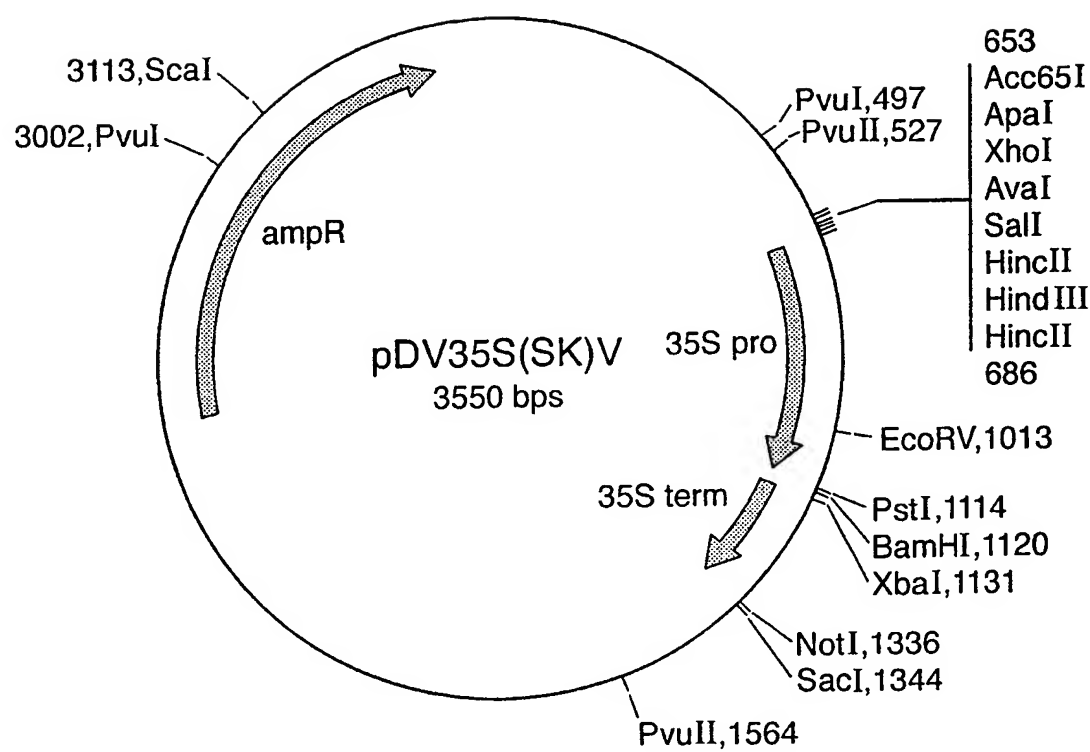


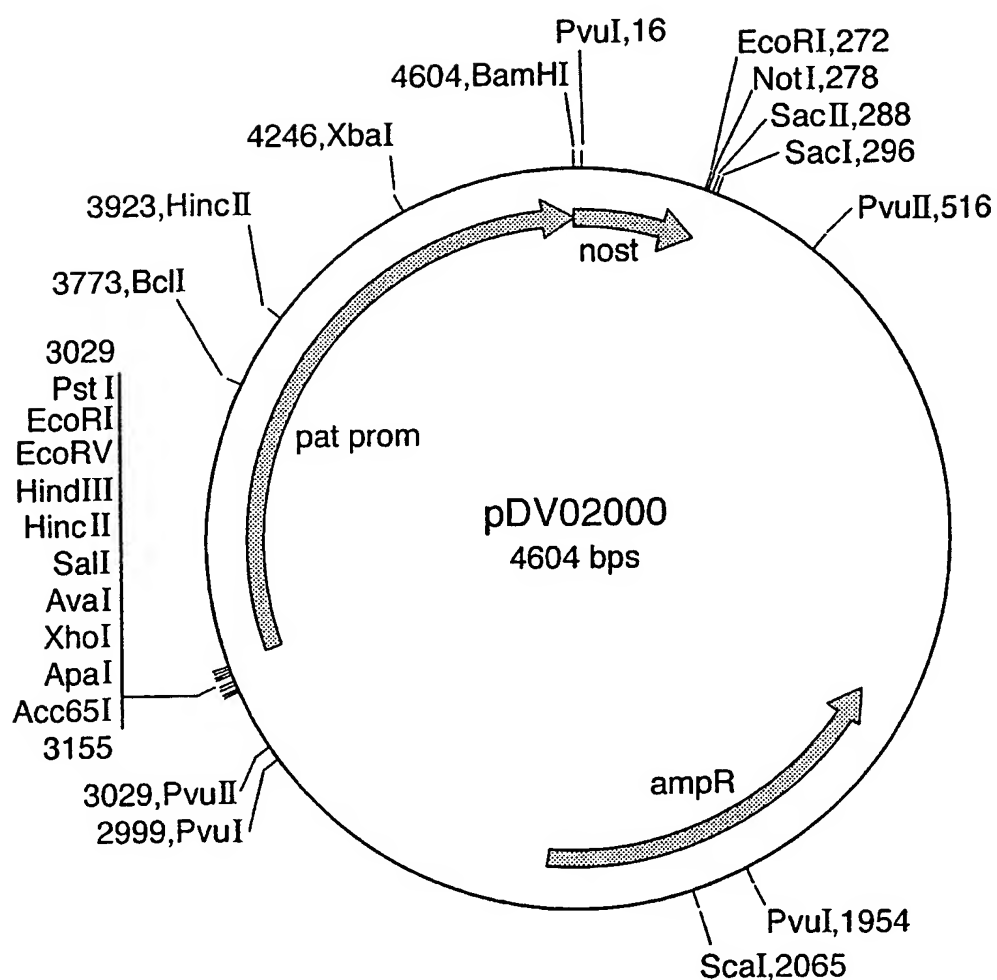
Fig.6.

Fig.7.

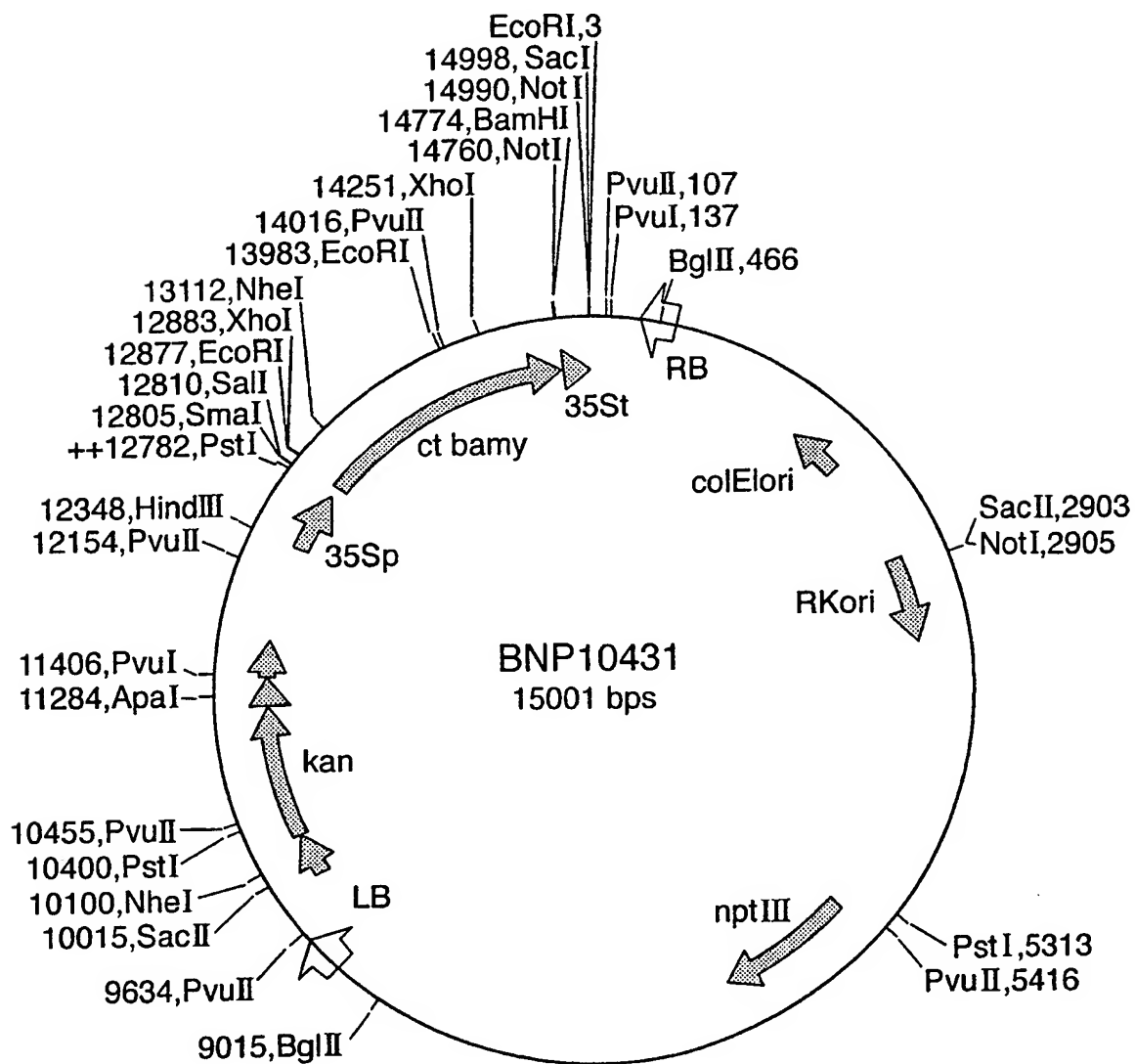


Fig.8.

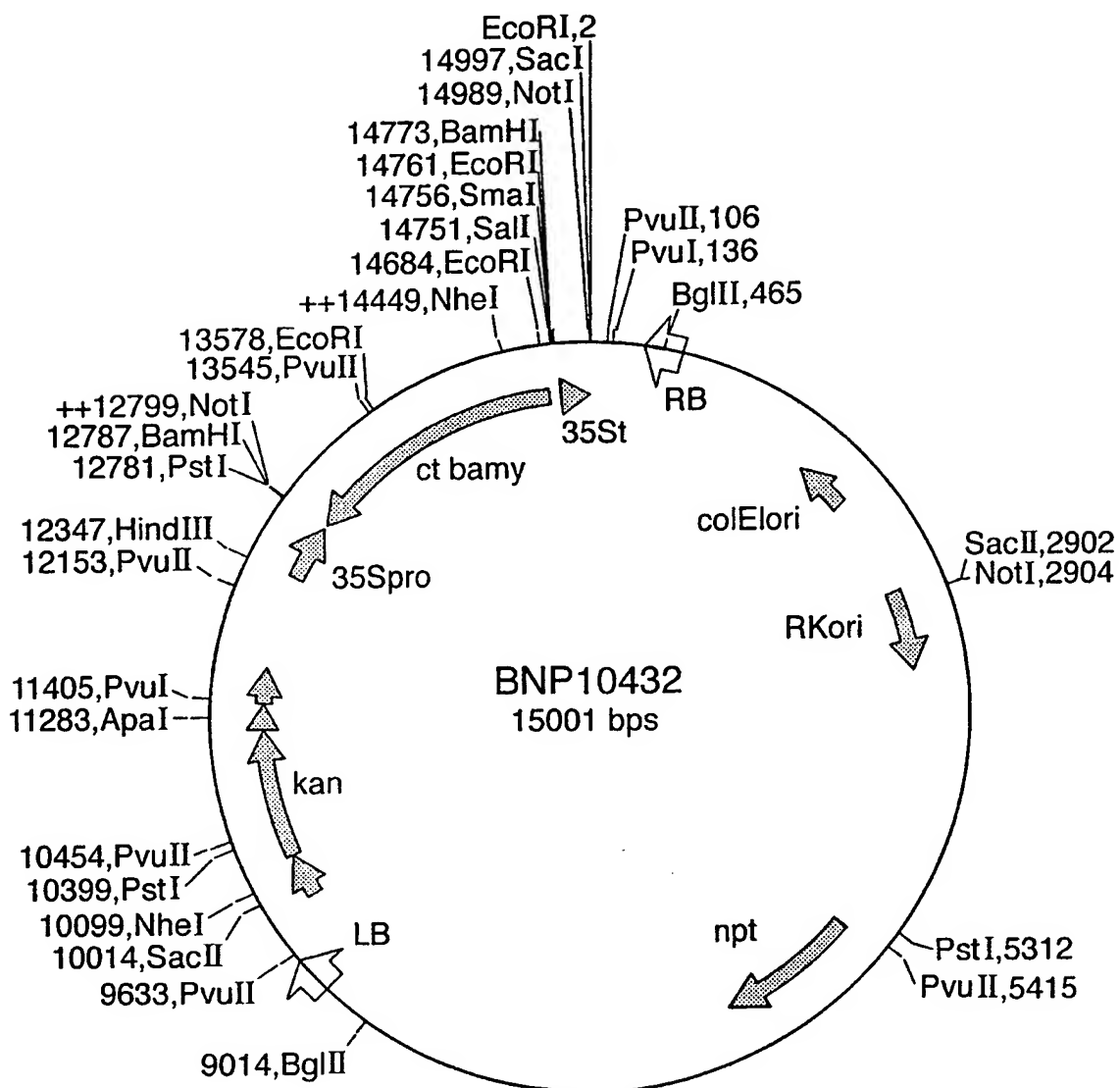
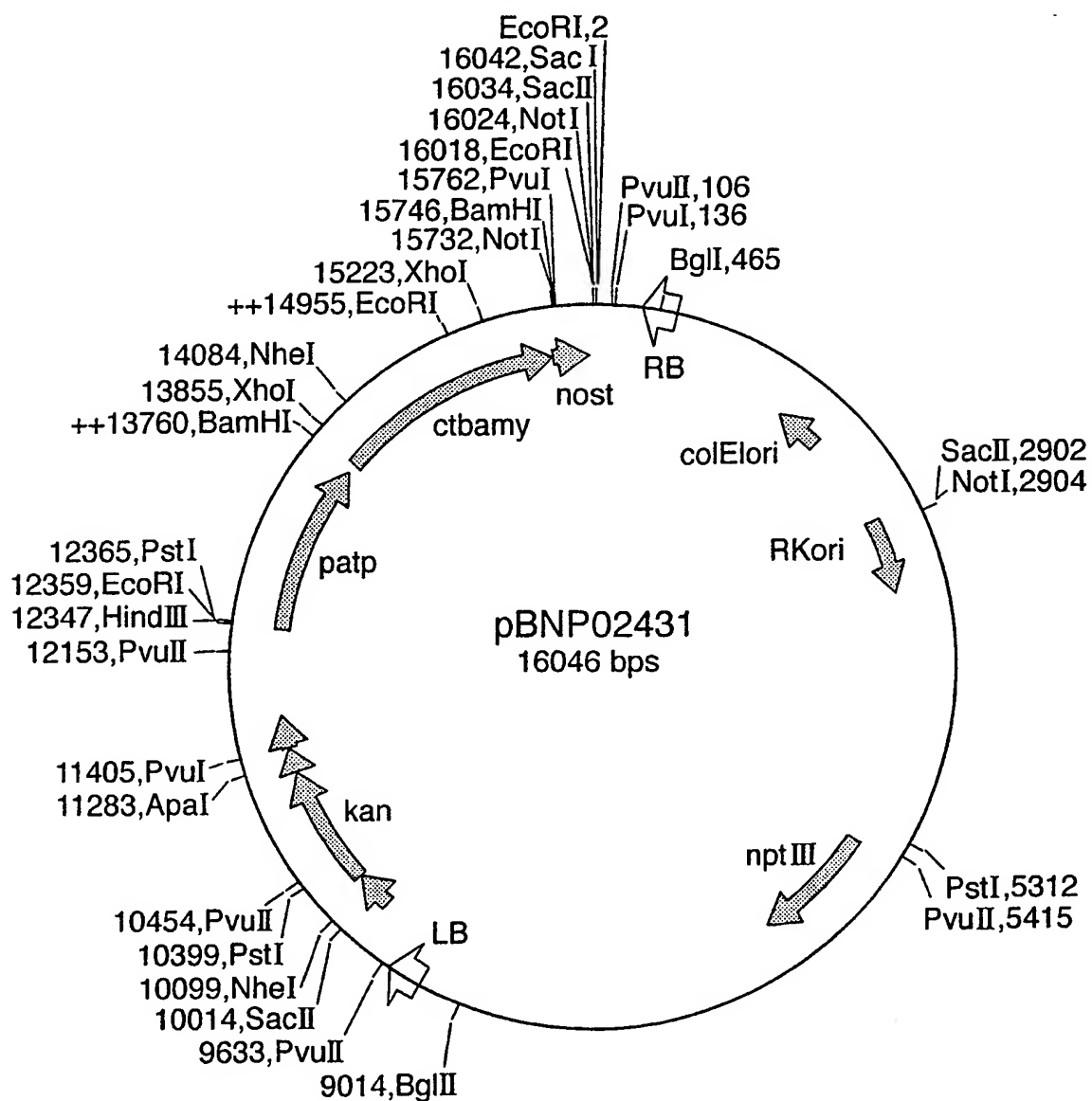
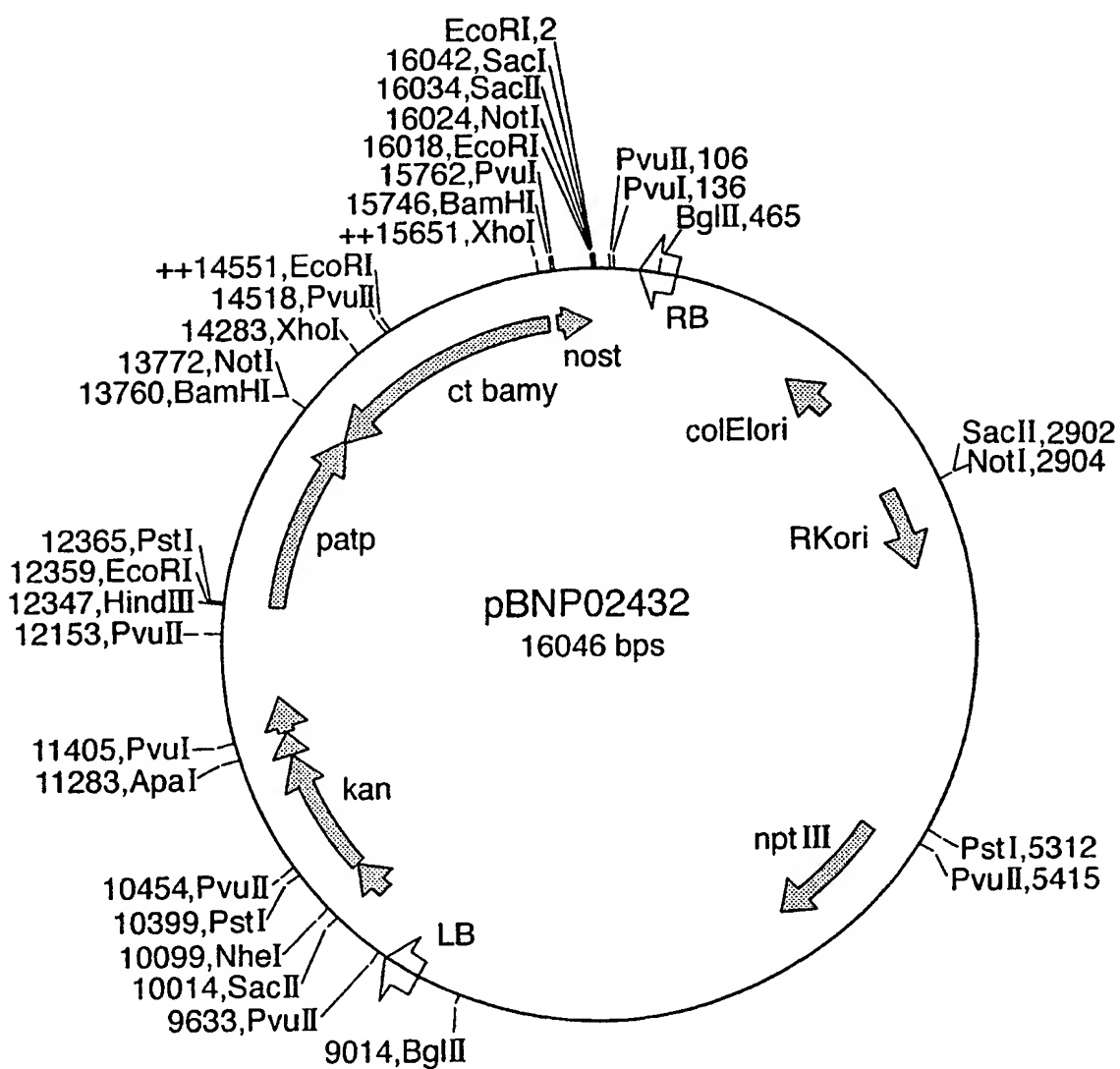


Fig.9.



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Fig.10.



SEQUENCE LISTING

(1) **GENERAL INFORMATION**

(i) **APPLICANTS:**

(A) NAME: Advanced Technologies (Cambridge) Limited
(B) STREET: Globe House, 1 Water Street
(C) CITY: London
(E) COUNTRY: England
(F) POSTAL CODE: WC2R 3LA

(ii) **TITLE OF INVENTION:** A novel plastid-targeting nucleic acid sequence, a novel β-amylase sequence, a stimulus-responsive promoter and uses thereof.

(iii) **NUMBER OF SEQUENCES:** 8

(iv) **CORRESPONDENCE ADDRESS:**

(A) ADDRESSEE: British American Tobacco (Investments) Limited
(B) STREET: Regents Park Road
(C) CITY: Southampton
(D) STATE: Hampshire
(E) COUNTRY: England
(F) POSTAL CODE: SO15 8TL

(v) **COMPUTER READABLE FORM:**

(A) MEDIUM TYPE: Diskette 3.50 inch
(B) COMPUTER: Viglen P5/75
(C) OPERATING SYSTEM: MS-DOS Windows 95
(D) SOFTWARE: Microsoft Word 97

(vi) **CURRENT APPLICATION DATA:**

(A) APPLICATION NUMBER: Not yet known
(C) CLASSIFICATION: Not yet known

(viii) **ATTORNEY/AGENT INFORMATION:**

(A) NAME: Mrs. M.R. Walford/ Mr. K.J.H. MacLean
(C) REFERENCE: RD-ATC-18

(ix) **TELECOMMUNICATION INFORMATION:**

(A) TELEPHONE: 01703 777155
(B) TELEFAX: 01703 779856

(2) INFORMATION FOR SEQ. ID. NO:1

- (i) SEQUENCE CHARACTERISTICS
- | | |
|-------------------|------------|
| (A) LENGTH: | 294 bps |
| (B) TYPE: | Nucleotide |
| (C) STRANDEDNESS: | Double |
| (D) TOPOLOGY: | Linear |
- (ii) MOLECULE TYPE: cDNA to mRNA
- (vi) ORIGINAL SOURCE:
- | | |
|------------------|----------------------|
| (A) ORGANISM: | Arabidopsis thaliana |
| (B) STRAIN: | Ecotype Colombia |
| (F) TISSUE TYPE: | Leaf |
- (vii) IMMEDIATE SOURCE:
- | | |
|--------------|-----------------------|
| (A) LIBRARY: | ABRC DNA Stock Centre |
| (B) CLONE: | EST 81E10T7 |
- (viii) POSITION IN GENOME:
- | | |
|-------------------------|---------------|
| (A) CHROMOSOME SEGMENT: | Chromosome IV |
|-------------------------|---------------|
- (ix) FEATURE:
- | | |
|---------------|-----------------|
| (A) NAME: | Transit peptide |
| (B) LOCATION: | 37-291 bp |
- (xi) SEQUENCE DESCRIPTION: SEQ. ID. NO:1:

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      S S L I K R K D A K S S R N Q E S S S N 28

121 AACATGACCTTGCGAAGATGAAGCCGCCAACATATCAGTTCCAAGCAAAGAACTCGGTT
      N M T F A K M K P P T Y Q F Q A K N S V 48

181 AAGGAAATGAAGTTCACTCACGGAGAAGACCTTCACGCCAGAAGGTGAAACCTTGAGAAA
      K E M K F T H E K T F T P E G E T L E K 68

241 TGGGAGAAGCTCCACGTTCTCATACCCACACTCCAAGAACGACGCTAGCGTT
      W E K L H V L S Y P H S K N D A S V           85

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(2) INFORMATION FOR SEQ. ID. NO:2

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1642
(B) TYPE: Nucleotide
(C) STRANDEDNESS: Double
(D) TOPOLOGY: Linear

(ii) MOLECULE TYPE: cDNA to mRNA

(vi) **ORIGINAL SOURCE:**
(A) ORGANISM: *Arabidopsis thaliana*
(B) STRAIN: Ecotype Columbia
(C) TISSUE TYPE: Leaf

(vii) **IMMEDIATE SOURCE:** ABRC DNA Stock Centre
(A) LIBRARY:
(B) CLONE: EST 81E10T7

(viii) **POSITION IN GENOME:** Chromosome IV
(A) **CHROMOSOME SEGMENT:**

(ix) FEATURE:	
(A) NAME:	Coding Sequence
(B) LOCATION:	1-1393bps
(D) OTHER INFORMATION:	Mature peptide

(xi) SEQUENCE DESCRIPTION: SEQ. ID. NO:2:

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      P R A M N A S L M A L K G A G V E G V M 40

121    GTGGATGCTTGGTGGGGATTGGTGGAGAAAGATGGACCTATGAATTATAACTGGGAAGGC
      V D A W W G L V E K D G P M N Y N W E G 60

181    TATGCCGAGCTTATACAGATGGTCAAAAGCACGGTCTCAAACCTCCAGGTCGTTATGTCA
      Y A E L I Q M V Q K H G L K L Q V V M S 80

241    TTCCATCAATGTGGAGGAAACGTTAGGAGACTCTTGCAGTATCCCCTGCCTCCATGGGTG
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301    CTTGAAGAGATCAGCAAGAACCCCTGATCTTGTCTACACAGACAAATCTGGAGAAGGAAC
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361    CCTGAATATATCTCCTGGGATGTGATTCTGTGCCTGCTTAAGAGGAAGAACACCTATC
      P E Y I S L G C D S V P V L R G R T P I 140

421    CAGGTCTACTCAGATTCTCATGAGGAGCTTCCGTGAACGATTGAAGGCTACATAGGAGGA
      O V Y S D F M R S F R E R F E G Y I G G 160

```

481 GTTATTGCGGAAATTCAAGTAGGAATGGGACCTTGTGGAGAATTGAGATACCCATCATA
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 541 CCTGAAAGCAACGGGACCTGGAGATTCCCCGGATTGGAGAGTTCCAGTGCTACGACAAG
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 601 TATATGAAATCGTCACTCAGGCATATGCTGAATCAATTGGAAAACTAACGGGGAAACA
 Y M K S S L Q A Y A E S I G K T N W G T 220

 661 AGCGGACCTCATGATGCCGGCGAGTACAAGAACCTCCCAGAAGATACTGAATTTCAGG
 S G P H D A G E Y K N L P E D T E F F R 240

 721 AGAGACGGAACATGGAATAGCGAGTATGAAAGTTTCATGGAATGGTACTCCGGGAAG
 R D G T W N S E Y G K F F M E W Y S G K 260

 781 CTGCTAGAACATGGAGACCAACTCCTATCTCAGCGAAAGGTATCTTCAAGGAAGCGGA
 L L E H G D Q L L S S A K G I F Q G S G 280

 841 GCAAAGCTATCAGGAAAGGTAGCTGGAATTCACTGGCACTACAACACCAGGTACACGCA
 A K L S G K V A G I H W H Y N T R S H A 300

 901 GCTGAGCTAACCGCTGGATATTACAACACAAGAAACCATGACGGTATCTGCCAATAGCT
 A E L T A G Y Y N T R N H D G Y L P I A 320

 961 AAGATGTTCAACAAACATGGAGTTGTGCTCAACTCACCTGCATGGAGATGAAAGACGGG
 K M F N K H G V V L N F T C M E M K D G 360

 1021 GAGCAACCTGAGCACCGAATTGCTCACAGAACGGTCTGGTCAAGCAAGTACAGAACCGG
 E Q P E H A N C S P E G L V K Q V Q N A 380

 1081 ACAAGGCAGGCCGGAACCGAACCTAGCAGGGAGAACCGCCTAGAACGATATGACTCGAGC
 T R Q A G T E L A G E N A L E R Y D S S 400

 1141 GCATTGGACAAGTGGTAGCAACAAATAGGTCAAGATTCTGAAATGGTTAACCGCATT
 A F G Q V V A T N R S D S G N G L T A F 420

 1201 ACTTACCTAAGAACAAAGCGTTATTGAGGGTCAAAATTGGCAGCAGTTAGTGGAG
 T Y L R M N K R L F E G Q N W Q Q L V E 440

 1261 TTTGTTAAGAACATGAAGGAAGGTGGTCACTGGAGGAGACTCTCAAAAGAACACAACT
 F V K N M K E G G H G R R L S K E D T T 460

 1321 GGAAGTGCACCTTATGTTGGATTGTCAAAGCAAGATCGCTGAGAATGTGGAGGAGGCT
 G S D L Y V G F V K G K I A E N V E E A 480

 1381 GCTTTAGTGTAAATTCCCACATAGGTACATACATATAGTGTGGTATTGTATTCCCTG
 A L V - 483

 1441 TCTGATAAAATAACTAGAGAGATCAAACCAAGTAAGAGTGTAAAGCTATAGATTTGCACAA

 1501 TTCTGGGTCAAGAGTCAGAGCAGAGCAAGAGCAAAATCAAGATGATGTACACTTAGATGTAT

 1561 CCTATGAGTTTCTTGTACATCATCTTACATCTTAATCTCAAATACTATGCATTTT

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(2) INFORMATION FOR SEQ. ID. NO:3

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH:	1953
(B) TYPE:	Nucleotide
(C) STRANDEDNESS:	Double
(D) TOPOLOGY:	Linear

(ii) MOLECULE TYPE: cDNA to mRNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM:	Arabidopsis thaliana
(B) STRAIN:	Ecotype Columbia
(F) TISSUE TYPE:	Leaf

(vii) IMMEDIATE SOURCE:

(A) LIBRARY:	ABRC DNA Stock Centre
(B) CLONE:	EST 81E10T7

(viii) POSITION IN CHROMOSOME:

(A) CHROMOSOME SEGMENT:	Chromosome IV
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(ix) FEATURE:

(A) NAME:	CDS
(B) LOCATION:	37-1683 bp
(C) OTHER INFORMATION:	Chloroplast targeted

(xi) SEQUENCE DESCRIPTION: SEQ. ID. NO:3:

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311     AACATGACCTTGCGAAGATGAAGCCGCCAACATATCAGTTCCAAGCAAAGAACTCGGTT
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461     AAGGAAATGAAGTTCACTCACGAGAAGACCTTCACGCCAGAAGGTGAAACCCTTGAGAAA
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611     TGGGAGAAGCTCCACGTTCTCTCATACCCACACTCCAAGAACGACGCTAGCGTTCCGGTG
                  W E K L H V L S Y P H S K N D A S V P V 88
761     TTCGTCATGTTACCGCTCGACACAGTAACAATGTCAGGGCATTGAAACAAACCACGAGCC
                  F V M L P L D T V T M S G H L N K P R A 108
911     ATGAACGCTAGTTGATGGCTCTGAAAGGAGCTGGTGTGGAAGGTGTGATGGTGGATGCT
                  M N A S L M A L K G A G V E G V M V D A 128
1061    TGGTGGGGATTGGTGGAGAAAGATGGACCTATGAATTATAACTGGGAAGGCTATGCCGAG
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```

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C G G N V G D S C S I P L P P W V L E E 188

601 ATCAGCAAGAACCCCTGATCTTGCTACACAGACAAATCTGGGAGAAGGAACCCCTGAATAT
I S K N P D L V Y T D K S G R R N P E Y 208

661 ATCTCCTTGGGATGTGATTCTGTGCCTGCTTAAGAGGAAGAACACCTATCCAGGTCTAC
I S L G C D S V P V L R G R T P I Q V Y 228

721 TCAGATTCATGAGGAGCTTCCGTGAACGATTGAAGGCTACATAGGAGGAGTTATTGCC
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781 GAAATTCAAGTAGGAATGGGACCTTGTGGAGATTGAGATAACCATCATACCCCTGAAAGC
E I Q V G M G P C G E L R Y P S Y P E S 268

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N G T W R F P G I G E F Q C Y D K Y M K 288

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T W N S E Y G K F F M E W Y S G K L L E 348

1081 CATGGAGACCAACTCCTATCTCAGCGAAAGGTATCTTCAAGGAAGCGGAGCAAAGCTA
H G D Q L L S S A K G I F Q G S G A K L 368

1141 TCAGGAAAGGTAGCTGGAATTCACTGGCACTACAACACCAGGTACACGCAGCTGAGCTA
S G K V A G I H W H Y N T R S H A A E L 388

1201 ACCGCTGGATATTACAACACAAGAACCATGACGGGTATCTGCCAATAGCTAAGATGTT
T A G Y Y N T R N H D G Y L P I A K M F 408

1261 AACAAACATGGAGITGTGCTCAACTTCACCTGCATGGAGATGAAAGACGGGAGCAACCT
N K H G V V L N F T C M E M K D G E Q P 428

1321 GAGCACCGAATTGCTCACAGAAGGTCTGGTCAAGCAAGTACAGAACCGCACAAGGCAG
E H A N C S P E G L V K Q V Q N A T R Q 448

1381 GCCGGAACCGAACTAGCAGGGAGAACCGCCTAGAACGATATGACTCGAGCGCATTGG
A G T E L A G E N A L E R Y D S S A F G 468

1441 CAAGTGGTAGCAACAAATAGGTCAAGATTCTGAAATGGGTAACCGCATTACTTACCTA
Q V V A T N R S D S G N G L T A F T Y L 488

1501 AGAATGAACAAGCGTTATTGAGGGTCAAAATTGGCAGCAGTTAGGGAGTTGTTAAG
R M N K R L F E G Q N W Q Q L V E F V K 508

1561 AACATGAAGGAAGGTGGTCATGGGAGGAGACTCTCAAAAGAACACAACTGGAAAGTGAC
N M K E G G H G R R L S K E D T T G S D 528

1621 CTTTATGTTGGATTGTCAAAGGCAAGATCGCTGAGAATGTGGAGGAGGCTGCTTAGTG
L Y V G F V K G K I A E N V E E A A L V 548

1681 TAATTCCCACATAGGTACATACATATAAGTGTGGTATTGTATTGCCTGTCTGATAAA

1741 TAACTAGAGAGATCAAACCAGTAAGAGTGTAAAGCTATAGATTGCACAATTCTGGGTC

1801 AGAGTCAGAGCAAAGAGAAGCAAATCAAGATGATGTACACTTAGATGTATCCTATGAGT

1861 TTTCCCTGTACATCATCTTCATACTCTTAATCTCAAATACTATGCATTTCTCCAAAAA

1921 AAAAAAAAAAGGGCGGCCGCTCTAGAGGATCC

(2) INFORMATION FOR SEQ. ID. NO:4

(i) SEQUENCE CHARACTERISTICS:

- | | |
|-------------------|-----------------|
| (A) LENGTH: | 21bp |
| (B) TYPE | Nucleotide |
| (C) STRANDEDNESS: | Single stranded |
| (D) TOPOLOGY: | Linear |

(xi) SEQUENCE DESCRIPTION: SEQ. ID. No. 4

AATTCCCTCGA GTTCTCTTAT C

21

(2) INFORMATION FOR SEQ. ID. NO:5

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21bp
(B) TYPE Nucleotide
(C) STRANDEDNESS: Single stranded
(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ. ID. No. 5

GGGGATCCCT GACATTGTTA C 21

2) INFORMATION FOR SEQ. ID. NO:6

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 35bp
(B) TYPE Nucleotide
(C) STRANDEDNESS: Single stranded
(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ. ID. No. 6

GCTGGTACTC CTGCAGGATC CGGTCCGGAA TTCCC 35

(2) INFORMATION FOR SEQ. ID. NO:7

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 35bp
(B) TYPE Nucleotide
(C) STRANDEDNESS: Single stranded
(D) TOPOLOGY: Linear

(xi) SEQUENCE DESCRIPTION: SEQ. ID. NO: 7

GGGAATTCCG GACCGGATCC TGCAGGCGTA CCAGC 35

SEQUENCE INFORMATION FOR SEQ. ID. NO. 8

(2) INFORMATION FOR SEQ. ID. NO:8

- (i) SEQUENCE CHARACTERISTICS:
- | | |
|-------------------|-----------------|
| (A) LENGTH: | 1652 bp |
| (B) TYPE | Nucleotide |
| (C) STRANDEDNESS: | Double stranded |
| (D) TOPOLOGY: | Linear |
- (ii) MOLECULE TYPE: Genomic DNA
- (vi) ORIGINAL SOURCE:
- | | |
|---------------|----------------------|
| (A) ORGANISM: | Arabidopsis thaliana |
| (B) STRAIN: | Ecotype Columbia |
| (C) TISSUE: | Leaf |
- (vii) IMMEDIATE SOURCE:
- | | |
|--------------|-------------------------------|
| (A) LIBRARY: | EU Arabidopsis Genome Project |
| (B) CLONE: | Cosmid G16599 |
- (viii) POSITION IN GENOME:
- | | |
|-----------------|--------------|
| (A) CHROMOSOME: | Chromosome 4 |
|-----------------|--------------|
- (xi) FEATURE:
- | | |
|---------------|---|
| (A) NAME: | Promoter |
| (B) LOCATION: | 17534-19185bp of G16599
Complementary strand |

(xi) SEQUENCE DESCRIPTION: SEQ. ID. NO: 8

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1 AAGCTTGTGT CTATTTCAAA TTCTTGACCG TAGATGTCAC AACATGCATA
51 TATCATTGAA AACAGAGCAA CACAGGAAAC CAAGCATATG TATCTAGATA
101 TACTTAGCAA GACATAACTA TAGTCTTGA ATCAACATAG GGATTAATGA
151 TAGAGAATGA GGAAGCTCAA GATTTTATAC TCAGTTCTT ACAAAACAAA
201 TTTCTCTCTA ACTGAAAAAA CACCAATTAG GATTGAAAGA GCGTACCTGT
251 TTGAGTCAAT GTCCAATGTC GTCCCCCCGC CTTCTACATT TCTTAGCCTG
301 CTGAATAAAA GCACAAGCCA AAATGAGAAG GTGCCAAAGG CGATAAGGAT
351 CAATTCTAC CATTAAAAAA ACTAATGGTG AGAATTAGAA ACGAGAGAAA
401 ACTACTTGTGTT GAGGAAATAG CCAAAAGCGC AATCTTCGTC ACCTGAATAA
451 AGACCAAACC GTCACTTTCA ATGAGTCAGC AAGAAAAAGA GAGAGAGAGA
501 GAGAGAGATT CTCTATAACA TTTGAGTCGA CATGGATTCT AATGCATCAA

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551 AAGTCATCTC CAATAAACAA ACACTTGAAA CTCACATGGC TAATAGAAC
601 AGATCAAAGC CTTAAGTATT AAGCATTACA GACACTACTG GCTAACTTTT
651 GACACATGTT CTTAAGTAAC ATAGTATCAA TATCCGTGAA TCACATCGAA
701 CACACACAAC AAGGGCTTAA TGCACTAAAG TCCTGTTATT TCCATATAAC
751 AACATATTTTC ATTTACAAAC AGAATGCAGC ATTCAAGGCAG TCCAAATGGA
801 AAGGTTGACA AAAAAATATA ATCTTGAAAC TCTACATATA TGGCAGAATG
851 TAATAACCAG GCAAGAAAAA AACAGAATAA ACAGATCAAT GAGTATGATA
901 TAAAAAAAAG TCACAAAGAA TGTGCCACAG TGAACAAGAG GGCCATGAGA
951 AGAAATTTTC AAAGAAAATA TTAGCATTGT TAGAATTTTT TGGGTCAATG
1001 GATCTGTCAG CTGCTTAGTT GGAAAACACA AATCTTACAG GAAGGAAAGT
1051 CCAAGAAAAA GAAAATAAGC AAAGTTAATA GCCACCACAA GAAATTTCAT
1101 ACAGAAAATAA TTAAATCGTT GCACTTATCT TCTTATTCAA ACTAAAATCA
1151 AGAGAACTTA ATAATTTCA GCCACACGAA CCATGTGTTCA AAAGCCAAG
1201 GTGAGAAGCC AAAATTATCA GCTTATCTCC ATTAACAAGG GAAAAGCAAG
1251 ACTAGATTAA AGAGTTCTCT GTAACTAAAA ACTGCAGGAG TGAGTAAGTA
1301 AATAATTACAC CAACAGGAAA ACAAAACTCA ATTATCTATA GCTGAATACA
1351 CATGTAAATG AGAATTTATT AACTAAAACA TCTTCCTTG TAACTGATGT
1401 GACATTACAA ATTTCATT TTGAGGTGTA AGAACCGTGT GACAAGTGAA
1451 AAGGTAAAAA TAAGCAACCT TTGTGATATT TTCTCTCCAC TTTTGAAAT
1501 TGGGTCTCCA ACCACAGCC AATCAATATT CTTTATAAAAT ACAAAACACAC
1551 AAACAGCATC TTTCTCTCAA ACACAAACAT ATCTTCTATC AAACACCAAC
1601 AGCTCTATTC TCTACCTCAT TTCTCATCAT AACAAAGAGA GAGAAAAAAA
1651 CT